

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶:

A61K 9/52, 47/30

A1

(11) International Publication Number: WO 98/32427

(43) International Publication Date: 30 July 1998 (30.07.98)

(21) International Application Number:

PCT/US98/01556

(22) International Filing Date:

27 January 1998 (27.01.98)

(30) Priority Data:

08/789,734

27 January 1997 (27.01.97)

US

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(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: THERAPEUTIC TREATMENT AND PREVENTION OF INFECTIONS WITH A BIOACTIVE MATERIAL ENCAPSULATED WITHIN A BIODEGRADABLE-BIOCOMPATIBLE POLYMERIC MATRIX

(57) Abstract

Novel burst-free, sustained release biocompatible and biodegradable microcapsules which can be programmed to release their active core for variable durations ranging from 1-100 days in an aqueous physiological environment. The microcapsules are comprised of a core of polypeptide or other biologically active agent encapsulated in a matrix of poly(lactide/glycolide) copolymer, which may contain a pharmaceutically acceptable adjuvant, as a blend of upcapped free carboxyl end group and end-capped forms ranging in ratios from 100/0 to 1/99.

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1	THERAPEUTIC TREATMENT AND
2	PREVENTION OF INFECTIONS WITH A BIOACTIVE MATERIALS
3	ENCAPSULATED WITHIN A BIODEGRADABLE-BIOCOMPATIBLE
4	POLYMERIC MATRIX
5	I. GOVERNMENT INTEREST
6	The invention described herein may be manufactured, used and licensed
7	by or for the Government for Governmental purposes without the payment to
8	use of any royalties thereon.
9	II. CROSS REFERENCE
10	This application is a continuation-in-part of U.S. Patent Application
11	Serial No. 08/590,973 filed January 24, 1996 which in turn is a
12	continuation-in-part of U.S. Patent Application Serial No. 08/446,149 filed
13	May 22, 1995, which in turn is a continuation of U.S. Patent Application Scient
14	No. 590,308 dated March 16, 1984.
15	Additionally, this application is a continuation-in-part of U.S. Patent
16	Application Serial No. 08/446,148 filed May 22, 1995, which in turn is a
17	continuation-in-part of U.S. Patent Application Serial No. 08/867,301 files
18	April 10, 1992 now U.S. Patent No. 5,417,986 issued May 23, 1995, which in
19	turn is a continuation-in-part of U.S. Patent Application Serial No. 590,388
50	filed March 16, 1984.

III. FIELD OF THE INVENTION

This invention relates to compositions comprising active core material(s) such as biologically active agent(s), drug(s) or substance(s) encapsulated within an end-capped or a blend of uncapped and end-capped biodegradable-biocompatable poly(lactide/glycolide) polymeric matrix useful for the effective prevention or treatment of bacterial, viral, fungal, or parasitic infections, and combinations thereof. In the areas of general and orthopedic surgery, and the treatment of patients with infectious or chronic disease conditions, this invention will be especially useful to physicians, dentists and veternarians.

IV. BACKGROUND OF THE INVENTION

Wounds characterized by the presence of infection, devitalized tissue, and foreign-body contaminants have high infection rates and are difficult to treat.

To prevent infection, in bone and soft tissue systemic antibiotics must be administered within 4 hours after wounding when circulation is optimal. This has been discussed by J.F. Burke in the article entitled "The Effective Period of Preventive Antibiotic Action in Experimental Incisions and Dermal Lesions", Surgery. Vol. 50, Page 161 (1961). If treatment of bacterial infections is delayed, a milieu for bacterial growth develops which results is complications associated with established infections. (G. Rodeheaver et al., "Proteolytic Enzymes as Adjuncts to Antibiotic Prophylaxis of Surgical Wounds", American Journal of Surgery. Vol. 127, Page 564 (1974)). Once infections are established it becomes difficult to systemically administer certain antibiotics for extended periods at levels that are safe and effective at the

1	wound site. Unless administered locally, drugs are distributed throughout the
2	body, and the amount of drug hitting its target is only a small part of the total
3	dose. This ineffective use of the drug is compounded in the trauma patient by
4	hypovolemic shock, which results in a decreased vascular flow to tissues. (L.
5	E. Gelin et al., "Trauma Workshop Report:Schockrheology and Oxygen
6	Transport", Journal Trauma. Vol. 10, Page 1078 (1970)).
7	Additionally, infections caused by multiple-antibiotic resistant bacterial
8	are on the up-swing and we are on the verge of a potential world-wide medical
9	disaster. According to the Centers for Disease Control, 13,300 patients died
10	in U.S. hospitals in 1992 from infections caused by antibiotic-resistant
11	bacteria. Methicillin-resistant S. aureus (MRSA) is rapidly emerging as the
12	"pathogen of the 90's":
13	a. Some major teaching hospitals in U.S. report that up to 40%
14	of strains of S. aureus isolated from patients are resistant to methicillin. Many
15	of these MRSA strains are susceptible only to a single antibiotic (vancomycin).
16	b. Should MRSA also develop resistance to vancomycin, the
17	mortality rate among patients who develop MRSA infections could approach
18	80%, thereby increasing the threat of this infectious killer.
19	Moreover, Vancomycin resistance is on the up-swing:
20	a. 20% of Enterococci are now resistant to vancomycin
21	b. In 1989, only one hospital in New York City reported
22	vancomycin-resistant Enterococci. By 1991, the number of hospitals reponing
23	vancomycin resistance rose to 38.

c. transfer of vancomycin-resistant gene (via plasmid) has been shown experimentally between Enterococcus and S. aureus.

Many major pharmaceutical companies around the world have either completely eliminated or significantly reduced their research and development programs in the area of antibiotic research. According to a 1994 report by the Rockefeller University Workshop in Multiple Antibiotic Resistant Bacteria, we are on the verge of a "medical disaster that would return physicians back to the pre-penicillin days when even small infections could turn lethal due to the lack of effective drugs."

Despite recent advances in antimicrobial therapy and improved surgical techniques, osteomyelitis (hard tissue or bone infection) is still a source of morbidity often necessitating lengthy hospitalization. The failure of patients with chronic osteomyelitis to respond uniformly to conventional treatment has prompted the search for more effective treatment modalities. Local antibiotic therapy with gentamicin-impregnated poly(methylmethacrylate) (PMMA) bead chains (SEPTOPAL TM, E. Merck, West Germany) has been utilized in Germany for the treatment of osteomyelitis for the past decade and has been reported to be efficacious in several clinical studies. The beads are implanted into the bone at the time of surgical intervention where they provide significantly higher concentrations of gentamicin than could otherwise be achieved via systemic administration. Serum gentamicin levels, on the other hand, remain extremely low thereby significantly reducing the potential for nephro- and ototoxicity that occurs in some patients receiving gentamicin systemically.

Administration for use in the United States, some orthopedic surgeons in this country are fabricating their own "physician-made beads" for the treatment of chronic osteomyelitis. A major disadvantage of the beads, however, is that because the PMMA is not biodegradable it represents a foreign body and should be removed at about 2-weeks postimplantation thereby necessitating in some cases an additional surgical procedure. A biodegradable-biocompitable, antibiotic carrier, on the other hand, would eliminate the need for this additional surgical procedure and may potentially reduce both the duration as well as the cost of hospitalization.

The concept of local, sustained release of antibiotics into infected bone is described in recent literature wherein antibiotic-impregnated PMMA macrobeads are used to treat chronic osteomyelitis. The technique as currently used involves mixing gentamicin with methylmethacrylate bone cement and molding the mixture into beads that are 7mm in diameter. These beads are then locally implanted in the infected site at the time of surgical debridement to serve as treatment. There are, however, significant problems with this method. These include: 1) initially, large amounts of antibiotics diffuse from the cement but with time the amount of antibiotic leaving the cement gradually decreases to subtherapeutic levels; 2) the bioactivity of the antibiotic gradually decreases; 3) methylmethacrylate has been shown to decrease the ability of polymorphonuclear leukocytes to phagocytize and kill bacteria; 4) the beads do not biodegrade and usually must be surgically removed; and 5) the exothermic reaction that occurs during curing of methymethacrylate limits the method to

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the incorporation of only thermostable antibiotics (primarily aminoglycosides).

Nevertheless, preliminary clinical trials using these beads indicate that they are equivalent in efficacy to longer term (4-6 weeks) administration of systemic antibiotics.

In many instances, infectious agents have their first contact with the host at a mucosal surface; therefore, mucosal protective immune mechanisms are of primary importance in preventing these agents from colonizing or penetrating the mucosal surface. Numerous studies have demonstrated that a protective mucosal immune response can best be initiated by introduction of the antigen at the mucosal surface, and parenteral immunization is not an effective method to induce mucosal immunity. Antigen taken up by the gutassociated lymphoid tissue (GALT), primarily by the Peyer's patches in mice, stimulates T helper cell (Th) to assist in IgA B cell responses or stimulates T suppressor cells (Ts) to mediate the unresponsiveness of oral tolerance. Particulate antigen appears to shift the response towards the (Th) whereas soluble antigens favor a response by the (Ts). Although studies have demonstrated that oral immunization does induce an intestinal mucosal immune response, large doses of antigen are usually required to achieve sufficient local concentrations in the Peyer's patches. Unprotected protein antigens may be degraded or may complex with secretory IgA in the intestinal lumen.

In the process of vaccination, medical science uses the body's innate ability to protect itself against invading agents by immunizing the body with antigens that will not cause the disease but will stimulate the formation of antibodies that will protect againts the disease. For example, dead organisms

are injected to protect against bacterial diseases such as typhoid fever and whooping cough, toxins are injected to protect against viral diseases such as 1 2 poliomyelitis and measles. It is not always possible, however, to stimulate antibody formation 3 merely by injecting the foreign agent. The vaccine preparation must be 4 immunogenic, that is, it must be able to induce an immune response. Certain 5 6 agents such as tetanus toxoid are innately immunogenic, and may be administered in vaccines without modification. Other important agents are not 7 immunogenic, however, and must be converted into immunogenic molecules 8 9 before they can induce an immune response. 10 The immune response is a complex series of reactions that can 11 generally be described as follows: 1. the antigen enters the body and encounters antigen-presenting cells 12 which process the antigen and retain fragments of the antigen on their surfaces; 13 2. the antigen fragment retained on the antigen presenting cells are 14 15 recognized by T cells that provide help to B cells; and 3. the B cells are stimulated to proliferate and divide into antibody 16 17 forming cells that secrete antibody against the antigen. Most antigens only elicit antibodies with assistance from the T cells 18 19 and, hence, are known as T-dependent (TD). These antigens, such as proteins, can be processed by antigen presenting cells and thus activate T cells 20 in the process described above. Examples of such T-dependent antigens are 21 22

tetanus and diphtheria toxoids.

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	Some antigens, such as polysaccharides, cannot be properly processed
1	and are not recognized by I cens. These and
2	do not require T cell assistance to elicit antibody formation but can activate B
3	do not require T cell assistance to the color of the cells directly and, hence, are known as T-independent antigens (TI). Such T-cells directly and, hence, are known as T-independent antigens (TI).
4	cells directly and, hence, are known as a steep independent antigens include H. influenzae type by polyribosyl-ribitol-phosphate
5	independent antigens include H. influenzae type of P
6	and pneumococcal capsular polysaccharides.
7	and pneumococcal capsular polyamers. T-dependent antigens vary from T-independent antigens in a number of
8	ways. Most notably, the antigens vary in their need for an adjuvant, a
9	nonspecifically enhance the immune response.
	antigens elicit only low level and antigens elicit only low level and antigens
10	about are administered with an adjuvant. At 15
11	diptheria, pertussis, tetalius, is
12	Insolubilization of TD antigens into an age
13	their immunogenicity, even in the absence
14	form can also enhance their transfer of the dy responses when
15	adjuvant. Golub ES and WO WES and Wo west and adjuvant. Golub ES and WO WES and west antibody responses when contrast, T-independent antigens can stimulate antibody response is generally of
16	contrast, T-independent antigens can be administered in the absence of an adjuvant, but the response is generally of
17	administered in the absence of an anjuvant,
18	lower magnitude and shorter duration. Four other differences between T-independent and T-dependent antigens
19	Four other differences between 1-independent
20	are:
21	are: a) T-dependent antigens can prime an immune response so that a
	memory response can be elicited upon secondary challenge with the same
22	responses are sumulated to a
23	antigen. Memory or secondary responses. attain significantly higher titers of antibody that are seen in primary responses.
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1	T-independent antigens are unable to prime the immune system for secondary
2	responsiveness.
3	b) The affinity of the antibody for antigen increases with time
4	after immunization with T-dependent but not T-independent antigens.
5	c) T-dependent antigens stimulate an immature or neonatal
6	immune system more effectively than T-independent antigens.
	d) T-dependent antigens usually stimulate IgM, IgGl, IgG2a, and
7 8	IgE antibodies, while T-independent antigens stimulate IgM, IgGl, IgG2b, and
9	IgG3 antibodies.
10	These characteristics of T-dependent vs. T-independent antigens provide
11	both distinct advantages and disadvantages in their use as effective vaccines.
12	T-dependent antigens can stimulate primary and secondary responses which are
13	long-lived in both adult and in neonatal immune systems, but must frequently
14	be administered with adjuvants. Thus, vaccines have been prepared using only
15	an antigen, such as diphtheria or tetanus toxoid, but such vaccines may require
16	the use of adjuvants, such as alum for stimulating optima responses.
	Adjuvants are often associated with toxicity and have been shown to
17	nonspecifically stimulate the immune system, thus inducing antibodies of
18	
19	specificities that may be undesirable. Another disadvantage associated with T-dependent antigens is that very
20	Another disadvantage associated with a component of the same and the same associated with a component of the component of the same associated with a component of the
21	small proteins such as peptides, are rarely immunogenic, even when
22	administered with adjuvants. This is especially unfortunate because many
23	synthetic peptides are available today that have been carefully synthesized to

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represent the primary antigenic determinants of various pathogens, and would 1 otherwise make very specific and highly effective vaccines. In contrast, T-independent antigens, such as polysaccharides, are able 2 to stimulate immune responses in the absence of adjuvants. Unfortunately, 3 however, such T-independent antigens cannot stimulate high level or prolonged 4 antibody responses. An even greater disadvantage is their inability to stimulate 5 an immature or B cell defective immune system (Mond J.J., Immunological 6 Reviews 64:99, 1982) Mosier DE, et al., J. Immunol. 119:1874, 1977). 7 Thus, the immune response to both T-independent and T-dependent antigens is 8 9 not satisfactory for many applications. With respect to T-independent antigens, it is critical to provide 10 protective immunity against such antigens to children, especially against 11 polysaccharides such as H. influenzae and S. pneumoniae. With respect to T-12 dependent antigens, it is critical to develop vaccines based on synthetic 13 peptides that represent the primary antigenic determinants of various 14 15 One approach to enhance the immune response to T-independent pathogens. 16 antigens involves conjugating polysaccharides such H. influenzae PRP (Cruse 17 18 19

One approach to enhance the immune response to 1-independent antigens involves conjugating polysaccharides such H. influenzae PRP (Cruse J.M., Lewis R.E. Jr. ed., Conjugate vaccines in Contributions to Microbiology and Immunology, vol. 10, 1989) or oligosaccharide antigens (Anderson PW, et al., J. Immunol. 142:2464, 1989) to a single T-dependent antigen such as tetanus or diphtheria toxoid. Recruitment of T cell help in this way has been shown to provide enhanced immunity to many infants that have been immunized. Unfortunately, only low level antibody titers are elicited, and

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only some infants respond to initial immunizations. Thus, several immunizationa are required and protective immunity is often delayed for months. Moreover, multiple visits to receive immunizations may also be difficult for families that live distant from medical facilities (especially in underdeveloped countries). Finally, babies less than 2 months of age may mount little or no antibody response even after repeated immunization.

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One possible approach to overcoming these problems is to homogeneously disperse the antigen of interest within the polymeric matrix of appropriately sized biodegradable-biocompatible microspheres that are specifically taken up by GALT. Eldridge et al. have used a murine model to show that orally-administered 1-10 micrometer microspheres consisting of polymerized lactide and glycolide, (the same materials used in resorable sutures), were readily taken up into Peyer's patches, and the 1-5 micrometer size were rapidly phagocytized by macrophages. Microspheres that were 5-10 micrometers (microns) remained in the Peyer's patch for up to 35 days, whereas those less than 5 micrometer disseminated to the mesenteric lymph node (MLN) and spleen within migrating MAC-1+ cells. Moreover, the levels of specific serum and secretory antibody to staphylococcal enterotoxin B toxoid and inactivated influenza A virus were enhanced and remained elevated longer in animals which were immunized orally with microencapsulated antigen as compared to animals which received equal doses of nonencapsulated antigen. These data indicate that microencapsulation of an antigen given orally may enhance the mucosal immune response against enteric pathogens. AF/R1 pili mediate the species-specific binding of E. coli RDEC-1

with mucosal glycoproteins in the small intestine of rabbits and are therefore an important virulence factor. Although AF/R1 pili are not essential for E. coli RDEC-1 to produce enteropathogenic disease, expression of AF/R1 to produce enteropathogenic disease, expression of AF/R1 promotes a more severe disease. Anti-AF/R1 antibodies have been shown to inhibit the attachment of RDEC-1 to the intestinal mucosa and prevent RDEC-1 disease in rabbits. The amino acid sequence of the AF/R1 pilin subunit has recently been determined, but specific antigenic determinants within AF/R1 have not been identified.

In the current study we have used these theortical criteria to predict probable T or B cell epitopes from the amino acid sequence of AF/R1. Four different 16 amino acid peptides that include the predicted epitopes have been synthesized: AF/R1 40-55 as a B cell epitope, 79-94 as a T cell epitope, 108-123 as a T and B cell epitope, and AF/R1 40-47/79-86 as a hybrid of the first eight amino acids from the predicted B cell epitope and the T cell epitope. We have used these peptides as well as the native protein to stimulate the in vitro proliferation of lymphocytes taken from the Peyer's patch, MLN, and spleen of rabbits which have received introduodenal priming with microencapsulated or non-encapsulated AF/R1. Our results demonstrate the microencapsulation of AF/R1 potentiates the cellular immune response at the level of the Peyer's patch, thus enhancing in vitro lymphocyte proliferation to both the native protein and its linear peptide antigens. CFA/I pili, rigid thread-like structures which are composed of repeating pilin subunits of 147 amino acid found on serogroups 015, 025, 078, and 0128 of enterotoxigenic E. coli (ETEC) (1-4,

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	the attachment to human brush bolders
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18). CF	A/I promotes mannose resistant attachment to human brush borders efore, a vaccine that established immunity against this protein may
	that established immunity against a
(5): ther	efore, a vaccine that control
(0)1	and subsequent disease. In addition,
rent	the attachment to host tissues and subsequent disease. In addition,
prevan	aning acid sequence homology
	the attachment to host ease. the CFA/I subunit shares N-terminal amino acid sequence homology the CFA/I subunit shares N-terminal amino acid sequence which
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	CEA/TV (CS4) (4), a substant
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Until recently, experiments to identify these epitopes were time consuming and costly; however, technology is now available which allows one to simultaneously identify all the T cell and B cell epitopes in the protein of interest. Multiple Peptide synthesis (Pepscan) is a technique for the simultaneous synthesis of hundreds of peptides on polyethylene rods (6). We have used this method to synthesize all the 140 possible overlapping actapeptides of the CFA/I protein. The peptides, still on the rods, can be used directly in ELISA assays to map B call epitopes (6. 12-14). We have also synthesized all the 138 possible overlapping decapeptides of the CFA/I protein. For analysis of T cell epitopes, these peptides can be cleaved from the rods and used in proliferation assays (15). Thus this technology allows efficient mapping and localization of both B cell and T cell epitopes to a resolution of a single amino acid (16). These studies were designed to identify antigenic epitopes of ETEC which may be employed in the construction of an effective subunit vaccine.

CFA/I pili consist of repeating pilin protein subunits found on several serogroups of enterotoxigenic E coli (ETEC) which promote attachment to

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human intestinal mucosa. We wished to identify areas within the CFA/I molecule that contain imunodominant T cell epitopes that are capable of stimulating the cell-mediated portion of the immune response in primates as well as immunodominant B cell epitopes. To do this, we (a) resolved the discrepancy in the literature on the complete amino acid sequence of CFA/I, (b) immunized three Rhesus monkeys with multiple i.m. injections of purified 5 6 CFA/I subunit in Freund's adjuvant, (c) synthesized 138 overlapping decapeptides which represented the entire CFA/I protein using the Pepscan 7 technique (Cambridge Research Biochemicals), (d) tested each of the peptides 8 for their ability to stimulate the spleen cells from the immunized monkeys in a 9 proliferative assay (e) synthesized 140 overlapping octapeptides which 10 represented the entire CFA/I protein, and (f) tested serum from each monkey 11 for its ability to recognize the octapeptides in a modified ELISA assay. A total 12 of 39 different CFA/I decapeptides supported a significant proliferative 13 response with the majority of the responses occurring within distinct regions of 14 the protein (peptides beginning with residues 8-40, 70-80, and 126-137). 15 Nineteen of the responsive peptides contained a serine residue at positions 2, 16 3, or 4 in the peptide, and a nine contained a serine specifically at position 3. 17 Most were predicted to be configured as an alpha holix and have a high 18 amphipathic index. Eight B cell epitopes were identified at positions 3-11, 11-19 21, 22-29, 32-40, 38-45, 66-74, 93-101, and 124-136. The epitope at position 20 11-21 was strongly recognized by all three individual monkeys, while the 21 epitopes at 93-101, 124-136, 66-74, and 22-29 were recognized by two of the 22 23 three monkeys. 24

Recent advances in the understanding of B cell and T cell epitopes have improved the ability to select probably linear epitopes from the amino acid sequence using theoretical criteria. B cell epitopes are often composed of a string of hydrophilic amino acids with a high flexibility index and a high probability of turns within the peptide structure. Prediction of T cell epitopes are based on the Rothbard method which identifies common sequence patterns that are common to known T cell epitopes or the method of Berzofsky and others which uses a correlation between algorithms predicting amphipathic helices and T cell epitopes.

V. SUMMARY OF THE INVENTION

This invention relates to active core materials such as biologically active agent(s), drug(s), or substance(s) encapsulated within a biodegradable-biocompatable polymeric matrix. In view of the enormous scope of this invention it will be presented herein as Phases I, II, and III. Phase I illustrates the encapsulation of antibiotics within a biodegradable-biocompatable polymeric matrix for the prevention and treatment of wound infections. Flase II illustrates the encapsulation of antigens (more specifically, oral-intestinal vaccine antigens) within a biodegradable-biocompatable polymeric matrix against diseases such as those caused by enteropathogenic organism. Phase III illustrates the use of a biodegradable-biocompatible polymeric matrix for barst-free programmable sustained release of biologically active agents, inclusive of peptides, over a period of up to 100 days in an aqueous physiological environment.

Controlled drug delivery from a biodegradable-biocompatable matrix offers profound advantages over conventional drug/antigen dosing. 1 Drugs/antigens can be used more effectively and efficiently, less drug/antigen 2 is required for optimal therapeutic effect and, in the case of drugs, toxic side 3 effects can be significantly, reduced or essentially eliminated through drug 4 targeting. The stability of some drugs/antigens can be improved allowing for a 5 longer shelf-life, and drugs/antigens with a short half-life can be protected 6 within the matrix from destruction, thereby ensuring sustained release of active 7 agent over time. The benefit of a continuous sustained release of drug/antigen 8 is beneficial because drug levels can be maintained within a constant 9 therapeutic range and antigen can be presented either continuously or in a 10 pulsatile mode as required to stimulate the optimal immune response. All of 11 this can be accomplished with a single dose of encapsulated drug/antigen. 12 This invention contemplates, but is not limited to, medically acceptable 13 methods for the effective local delivery of biologically active agents that, of 14 themselves, are directly (e.g. drugs, such as antibiotics) or indirectly (e.g. 15 vaccine antigens) therapeutic or prophylactic. It also includes drugs/agents that 16 17 elicit/modulate natural biological activity. Wounds characterized by the presence of infection, devitalized tissue, 18 and foreign-body contaminants have high infection rates and are difficult to 19 treat. This invention describes antibiotic formulation encapsulated within 20 microspheres of a biodegradable-biocompatable polymer that, when applied 21 locally to contaminated or infected wounds, provides immediate, direct, and 22 sustained (over a period up to 100 days), high concentrations of antibiotic is 23

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	the wound site (soft tissue and bone). By encapsulating antibiotics and
1	applying them directly, one can achieve a significant reduction in nonspecific
2	applying them directly, one can active a phenomena commonly observed binding of the drug to body proteins, a phenomena commonly observed
3	following conventional systemic administration of free drugs. Thus, less drug
4	following conventional systemic administration of the site of need, and
5	is required, higher concentrations are maintained at the site of need, and
6	efficacy is enhanced. This approach provides superior treatment over
7	conventional systemic administration of antibiotics for wound infections
8	because higher bacteriocidal concentrations can be achieved and maintained in
9	the wound environment. Higher concentrations kill more bacteria.
10	Applicants' invention for this application is described in Phase I.
11	Applicants' invention for this applicants reasoned that a protective mucosal immune response Furthermore, applicants reasoned that a protective mucosal immune response
12	Furthermore, applicants reasonable might be best initiated by introduction of an antigen at the mucosal surface,
13	because unprotected protein antigens delivered in a free form may be degraded
14	and with secretory IgA in the intestinal function production
15	processing in local immune cells. The results
16	antigen small enough in size to be pring
17	isioned as being able to induce and
	a plicants' invention for this application is
18	applicants propose using sover
19	including: 1) the direct application of the
20	application of drugs including. 2) oral delivery that provides either local drug to a surgical/traumatized area, 2) oral delivery that provides either local
21	drug to a surgical/traumatized artigen/drugs at mucosal membranes or deposition of microencapsulated antigen/drugs at mucosal membranes or
22	deposition of microencapsulated and arrayide local adherence of
23	transport across these membranes to provide local adherence of
24	microencapsulated drugs/antigen to mucosal membranes to provide sustained

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release of drug/antigen into soft tissue or a body cavity, and/or 3) sustained intercellular or extracellular drug/antigen release following subcutaneous injection.

In those instances where antibiotics are administered locally, applicants have found that the controlled release of the antibiotic from within a biodegradable-biocompatable polymeric matrix within 14 days to about 4 weeks without significant drug trailing is especially useful. However, if desired, the release of a biologically active agent from a polymeric matrix comprised of an active agent and a blend of uncapped and end-capped biodegradable poly DL(lactide-co-glycolide), can be controlled over a period of 1 to about 100 days without significant drug dumping or trailing. Such novel biocompatible-biodegradable microspheres developed with a burst-free programmable sustained release of biologically active agents, inclusive of polypeptides, are described in applicants' U.S. Patent Application Serial No. 08/590,973 filed January 24, 1996.

When antibiotics are administered systemically in the conventional manner, or locally as contemplated by the applicants, the immune response to the antibiotic and the potential for hypersensitivity and/or anaphylactoid response (especially to beta-lactam antibiotics such as penicillins/ampicillins) is a clinical concern. In early studies the inventors observed a specific IgG response to ampicillin as it was released from the microencapsulated formulation (illustrated in the histogram, Figure 1 and 2). This response is reminiscent of antibody elicited by vaccine antigens in conventional vaccines. The response to vaccine antigens is known to be accentuated by the use of an antigens in conventional vaccines.

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adjuvant such as alum. Alum is a c	rude, less adaptable delivery vehicle than		
its counterpart, the biodegradable-bi	ocompatable poly DL(lactide-co-glycolide),		
of this invention - the polymeric ma	trix. This knowledge stimulated additional		
studies relevant to the effects of sustain release of agents on the immune			
response.			

There are, in general, two forms of localized delivery which can be achieved with PLGA microspheres-delivery which is localized to individual cells of the body (intracellular delivery); and delivery which is localized to tissues within a specific region of the body (localized extracellular delivery).

Applicants have prepared antibiotic and hepatitis vaccine formulations which functioned by delivering localized extracellular doses of their active agents. This was achieved by using relatively large microspheres which served as a depot for the drug or antigen. Their large size 40-100 microns in diameter precluded their being phagocytized or diffusing throughout the intercellular fluid compartments of the body. Their drug agent loads were thus released within their immediate vicinity which resulted in the generation of very high local concentrations of antibiotic or the release of sufficiently high concentrations of free antigen to induce an immune response.

The large-diameter antibiotic bearing microspheres were originally developed by applicants primarily for topical application on exposed debrided tissues of combat wounds. However, an inherent property exhibited by the antibiotics when topically applied to a wound site is the generation of measurable levels of immune response. This concept of local delivery by

of therapeutic agents was subsequently applied to the development of an oral vaccine for protection against traveler's diarrhea caused by E. coli. Vaccine antigen was encapsulated into microspheres whose diameters were predominantly in the 5-10 micron size range based on an understanding that microspheres of this size would not readily be either phagocytized or transported across the gut wall into the body. Ingestion of these microspheres thus constituted a localized delivery achieved by topical application of the spheres to the wall tissue of the gut. This topical application resulted in the localized trapping of a small percentage of these sphere into the Peyer's patches where the spheres proceeded to release their antigen in a localized fashion to immune cells located within the intestinal Patches.

The concept of localized sustained local delivery has been further extended to the delivery of analgesics and anesthetics to exposed dental pulp to control pain and inflammatory responses. Again, the PLGA microsphere used for this type of delivery are relatively large (40-100 um in diameter) and serve as a topical depot for localized extracellular release of the drug.

Properties exhibited by active core materials in vivo, applicants have movel on to other non-topical application methods of using their microsphere delivery system. Some of these center on the use of small diameter microspheres ranging from sub micron to under 5 microns in diameter. These spheres show intracellular targeting of drug or antigen. They also allow for transmucosidelivery of drugs or antigens. The concept of localized delivery in these

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instances refers to the localized delivery of drug or agent within individual target cells of the body regardless of their location or distribution within the body. This approach is useful in development of antitubercular, antimalarial, antiviral, and antichlamydial formulations against intracellular parasites. It is also useful for the development of vaccines against intracellular parasites and for direct delivery of agents to presenting cells of the immune system.

Another nontopical application method of using PLGA microspheres resides in their usefulness as injectable depots for drugs intended for either localized or systemic delivery. Typically larger diameter microspheres are used for depots as these are less likely to diffuse away. The local or systemic nature of these delivery systems is, in part a function of the release rate of the drug from the depot and the diffusional and solubility characteristics of the drug being released. Cancer chemotherapeutics, systemic antibiotics, delivery of antibiotics to infected bone are potential application of this system.

Additional this non-topical systemic depot application can be extended to the injection of cancer-agent laden microspheres to embolize and destroy a malignant tumor. Additionally, the PLGA microspheres can be used as a carrier to deliversubstances useful for the in modification of cells or genes in bioengineering or genetic procedures.

Interest in the concept that antigens encapsulated within a biodegradable-biocompatible polymeric matrix could be formulated as a vaccine with superior efficacy over conventional vaccines, originated from the inventors' own observations that the drug, ampicillin, when sustain release from poly DL(lactide-co-glycolide) elicited antibody production. In these

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studies, the applicants were able to measure specific IgG antibodies to free ampicillin and to ampicillin released from microencapsulated ampicillin formulations in the sera of mice previously "treated" with the ampicillin formulations using ELISA. Numerous other studies also document the ability of beta-lactam antibiotic to elicit antibody. Selected, more recent studies whose findings are consistent with earlier discoveries made by applicants when conducting experiements with ampicillin include those by Klein et al. (1993) who detected specific IgG antibodies (IgG and IgG3 subclasses) to the Blactam ring in patients receiving penicillin therapy, work by Nagakura et al. (1990) which detected specific antibodies to cephalexin, a B-lactam antibiotic in the sera of guinea pigs, and Auci et al. (1993) who detected benzyl penicilloyl specific IgM, IgG IgE, and IgA antibody forming cells in lymphoid cells of mice given benzyl penicilloyl-Keyhole Limpet Hemocyanin. Pharmaceutical compositions of antigens encapsulated with poly DL(lactide-coglycolide) are described in Phase II. The microspheres of the invention allow for introduction of vaccine antigens to mucosal surfaces in particles that can be subsequently taken up locally by phagocytic cells. Such an approach for both drugs and antigens provides significant advantages in potency and efficacy over conventional systemically administered drugs or vaccines. A partial list of biologically active agents or drugs that will potentially derive significant medical benefits from this delivery system includes: antibacterial agents; peptides; polypeptides; antibacterial peptides; antimycobacterial agents; antimycotic agents; antiviral agents; antiparastic agents;, antifungal; antiyeast agents; hormonal peptides; cardiovascular agents; hormonal

peptides; cardiovascular agents; narcotic antagonists; analgesics; anesthetics; 1 insulins; steroids including HIV therapeutic drugs (including protease inhibitors) and AZT; estrogens; progestins; gastrointestinal therapeutic agents; 2 3 non-steroidal anti-inflammatory agents; parasympathoimetic agents; psychotherapeutic agents; tranquilizers; decongestants; sedative-hypnotics; non-4 estrogenic and non-progestional steroids; sympathomimetic agents; vaccines; 5 6 vitamins; nutrients; anti-migraine drugs; electrolyte replacements; ergot 7 alkaloids; anti-inflammary agents; prostaglandins; cytotoxic drugs; antigens; 8 antibodies; enzymes; growth factors; immunomodulators; pheromones; 9 prodrugs; psychotropic drugs; nicotine; antiblood clotting drugs; appetite 10 suppressants/stimulants and combinations thereof; contraceptive agents include estrogens such as diethyl silbestrol; 17-beta-estradiol; estrone; ethinyl estradiol; 11 mestranol; progestins such as norethindrone; norgestryl; ethynodiol diacetate; 12 lynestrenol; medroxyprogesterone acetate; dimethisterone; megestrol acetate; 13 14 chlormadinone acetate; norgestimate; norethisterone; ethisterone; melentate, 15 norgestimate; norethisterone; ethisterone; melengestrol; norethynodrel; and 16 spermicidal compounds such as nonyphenoxypolyoxyethylene glycol; benzethonium chloride; chlorindanol; include gastrointestinal therapeutic agents 17 18 such as aluminum hydroxide; calcium carbonate; magnesium carbonate; 19 sodium carbonate and the like; non-steroidal antifertility agents; 20 parasympathomimetic agents; psychotherapeutic agents; major tranquilizers 21 such as chloropromaquine HCL; clozapine; mesoridazine; metiapine; 22 reserpine; thioridazine; minor tranquilizers such as chlordiazepoxide; diazepam; meprobamate; temazepam and the like; rhinological decongestant; 23 24

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	sedative-hypnotics such as codeine; phenobarbital; sodium pentobarbital;
1	the steroids such as testosterone and testosterone
2	propionate; sulfonmides; sympathomimetic agents; vaccines; vitamins and
3	nutrient such as the essential amino acids; essential fats; anti-HIV agents;
4	nutrient such as the essential animo access animoquinolines; 8 aminoquinolines; including AZT; antimalarials such as 4-aminoquinolines; 8 aminoquinolines;
5	including AZT; antimalarials such as mazindol; phentermine; anti-
6	pyrimethamine; anti-migraine agents such as mazindol; phentermine; anti-
7	Parkinson agents such as L-dopa; antispasmodics such as atropine;
8	Parkinson agents such as Boop methscopolamine bromide; antispasmodics and anticholingeric agents such as
0	linestents: enzymes and the like; antitussives such as
9	noscapine: bronchodilators; cardiovascular agent
10	dextromethorphan and noscapatory as anti-hypertensive compounds; Rauwolfia alkaloids; coronary vasodilators;
11	as anti-hypertensive compounds, 222
12	as anti-hypertensive companies anti-hypertensive companies; pentaerythriotetranitrate; electrolyte nitroglycerin; organic nitrites; pentaerythriotetranitrate; electrolyte
	the potacsium chloride; ergotalkaloids such an organization
13	hydrogenated ergot alkaloids, universes govern
14	disadesergocomine methanesulfonate, university
15	combinations thereof; alkaloids such as appear
16	methaneusulfate and comonates methaneusulfate and comonates Belladonna; hyoscine hydrobromide; analgesics; narcotics such as codeine; Belladonna; hyoscine hydrobromide; analgesics; narcotics such as salicylates;
17	Belladonna; hyoscine hydrootomae, dihydrocodienone; meperidine; morphine; non-narcotics such as salicylates;
18	dihydrocodienone; meperidine, morphare; antibiotics such as the
19	dihydrocodienone, inspersor di
20	aspirin; acetaminophen; and a property containing ceflacor and cefuroxime; chloranphenical; gentamicin; cephalosporins including ceflacor and cefuroxime; chloranphenical; gentamicin;
21	cephalosporins including of Kanamycin A. Kanamycin B; the penicillins; ampicillin; amoxicillin;
22	A: chloropamtheniol; metromicatory
23	renicillin G; the tetracyclines; including inmices
	oxytetracycline periodized periodized oxytetracycline periodized periodized quinolones including ciprofloxacin; ofoxacin; macrolides including
24	dimination -

	clarithromycin; frythromycin; aminioglycosides including gentamicin;
1	clarithromycin; frythromycin; animogey amikacin; tobramycin and kanamycin; beta-lactams including ampacillin;
2	amikacin; tobramycin and kanamycin, ocu anamycin; tobramphenicol; fusidans;
3	polymyxin-B; amphotercin-B; aztrofonam; chloramphenicol; fusidans;
4	polymyxin-B; amphotered =, lincosamides; metronidazole; nitro-furantion; imipenem/cilastin; quinolones; lincosamides; trimethoprim;
5	systemic antibodies including rifampin; polygenes; sulfunamides; trimethoprim;
6	glycopeptides including vancomycin; teicoplanin and imidazoles; anti-cancer
7	glycopeptides including validations, anti-convulsants such as mephenytoin; agents; including anti-kaposi's sarcoma; anti-convulsants such as mephenytoin;
8	anti-emetics such as themsiperature,
9	as chlorophinazine; dimenhydrinate; dipliciniyas
	and the like; anti-inflammatory agents been
10	hydrocortisone; prednisolone; prednisole, non men
11	and the suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims water-soluble hormone drugs, and see a suringly for claims and see a suringly for
12	infolmmatory agents; antipyrencs, analgester,
13	sedatives; muscle relaxants; anucphicpass,
14	antiallergic drugs; cardiotonics; antiarriyamen
15	agents; antidepressants, antimore, anticoagulants; and antinarcotics; in vasodilators; antihypertensives; diuretics; anticoagulants; and antinarcotics; in
16	of 100-100,000 daltons; indoniculation,
17	the molecular wight range of 100 100 100; phenylbutazone; prostaglandins; cytotoxic drugs such as thiotepa; chloramucil; phenylbutazone; prostaglandins; cytotoxic drugs such as thiotepa; chloramucil;
18	phenylbutazone; prostaglandins; cywioxio and phenylbutazone; prostaglandins; cywioxio and cyclophosphamide; melphala; nitrogen mustard; methotrexate; antigens such as cyclophosphamide; melphala; nitrogen mustard; methotrexate; synthetic
19	cyclophosphamide; melphala; nitrogen meetro,
20	proteins; glycoproteins; synthetic peptides; carbohydrates; synthetic
21	proteins; glycoproteins, synthetic polysaccharides; lipids; glycolipids; lipopolysaccharides(LPS); synthetic
22	polysaccharides; lipids, growing attached adjuvants such as synthetic lipopolysaccharides and with or without attached adjuvants such as synthetic
23	designitives, antigens of such microoliganisms
24	muramyl dipeptide derivativos, estas gonorrhea; Mycobacterium tuberculosis; Picarinii Pnfumonia; Herpes virus
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(humonis types 1 and 2); Herpes zoster; Candidia albicans; Candida tropicalis; 1 Trichomonas vaginalis; Haemophilus vaginalis; Group B streptoccoccus ecoli; 2 Microplasma hominis; Hemophilus ducreyi; Granuloma inguimale; 3 Lymphopathia venerum; Treponema palidum; Brucela aborus Brucela meitensis 4 Brucela suis; Brucella canis Campylobacter fetus; Campylobacer fetus 5 intesinalis; Leptospira pomona. Listeria monocytogenes; Brucella ovis; Equine 6 herpes virus 1; Equine arteritis virus; IBR-IBP virus; Chlamydia psittaci; 7 Trichomonas foetus; Taxoplasma gondii; Escherichia coli; Actinobacillus 8 equuli; Salmonella abortus ovis. Salmonella abortus eui; Pseudomonas 9 aeruginosa; Corynebacterium equi; Corynebacterium pyogenes; Actinobaccilus 10 seminis; Mycoplasma bovigenitalium; Aspergilus fumigatus; Absidia ramosa; 11 Trypanosoma equiperdum; Babesia cabali; Clostridium tetani; antibodies which 12 counteract the above microorganisms; and enzymes such as ribonuclease; 13 neuramidinase; trypsin; glycogen phosphorylase; sperm lactic dehydrogenase; 14 sperm hyaluronidase; adenossinetriphosphase; alkaline phosphatase; alkaline 15 phospha esterase; amino peptides; typsin chymotrypsin amylase; muramidase; 16 acrosomal proteinase; diesterase; glutamic acid dehydrogense; succunic and 17 dehydrogenase; beta-glycophosphatase lipase; ATP-ase alpha-peptate gamma-18 glutamyiotranspeptidase; sterold-beta-ol-dehydrogenase; DPN-di-aprorase; and 19 combinations thereof. Having generally described the invention; a further 20 understanding can be obtained by reference to certain specific examples which 21 are provided herein for purpose of illustration only and are not intended to be 22 limiting unless otherwise specified. Moreover; the polymeric matrix of this 23 invention may be used for the in situ production and controlled release of 24

Additionally; effective testing or monitoring devices for chemical agents or bioactive agents can be made by encapsulating reagents which react as they are released from the polymeric matrix, with agents sought to be detected. The novel delivery system of this invention is applicable to all categories of active substances capable of being used for the prevention and/or treatment of human, animal and plant diseases. This delivery system is also applicable to the design of novel diagnostic tests. Additionally, it can be useful for the delivery to a subject of a polyfunctional mixture or cocktail formulation of encapsulated active (i.e. biologically) substances for the prevention and/or treatment of diseases the same or different. The encapsulated formulation ingredients would be comprised of multiple drugs, multipe vaccines or a combination thereof.

Applicants contemplate that the invention will be useful in the formulation of disease specific compositions for the prevention and/or treatment of diseases and/or ailments which include: viral infections; bacterial infections; fungal infections; yeast infections; parastic infections and more specific diseases and/or ailments; such as as, aids; alzheimer's dementia; angiogenesis diseases; aphthour ulcers in AIDS patients; asthma; atopic dermatitis; psoriasis; basal cell carcinoma; benign prostatic hypertrophy; blood substitute, blood substitute in surgery patients; blood substitute in trauma patients; breast cancer; breast cancer; cutaneous & metastatic; cachexia in AIDS; campylobacter infection; cancer; pnemonia; sexually transmitted diseases (STDs); cancer; viral dieases; candida albicians in AIDS and cancer; candidiasis in HIV infection; pain in

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cancer; pancreatic cancer; parkinson's disease; peritumoral brain edema; postoperative adhesions (prevent); proliferative diseases; prostate cancer; ragweed allergy; renal disease; restenosis; rheumatoid arthritis; rheumatoid arthritis; allergies; rotavirus infection; scalp psoriasis; septic shock; small-cell lung cancer; solid tumors; stroke; thrombosis; type I diabetes; type I diabetes w/kidney transplants; type II diabetes; viseral leishmaniasis; malaria; periodontal or gum disease; cardiac rthythm disorders; central nervous system diseases; central nervous system disorders; cervical dystonia (spasmodic torticollis); choridal neovascularization; chronic hepatitis c, b and a; colitis associated with antibiotics; colorectal cancer; coronary artery thrombosis; cryptosporidiosis in AIDS; cryptosporidium parvum diarrhea in AIDS; cystic fibrosis; cytomegalovirus disease; depression; social phobias; panic disorder; diabetic complications; disabetic eye disease; diarrhea associated with antibiotics; erectile dysfunction; genital herpes; graft-vs host disease in transplant patients; growth hormone deficiency; head and neck cancer; head 14 trauma; stroke; heparin neutralization after cardiac bypass; hepatocellular 15 carcinoma; HIV; HIV infection; huntington's disease; CNS diseases; 16 hypercholesterolemia; hypertension; inflammation; inflammation and 17 angiogensis; inflammation in cardiopulmonary bypass; influenza; migrain head 18 ache; interstitial cystitis; kaposi's sarcoma; kaposi's sarcoma in AIDS; lung 19 cancer; melanoma; molluscum contagiosum in AIDS; multiple sclerosis; 20 neoplastic meningitis from solid tumors; non-small cell lung cancer; organ 21 transplant rejection; osteoarthritis; rheumatoid arthritis; osteoporosis; drug 22 23 addiction; shock; ovarian cancer; and pain. 24

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Also contemplated here are those diseases or health conditions capable being benefitted by the list of biologically active agents or drugs previously 1 2 listed in the Summary of the Invention. 3

EFFECTS OF MICROENCAPSULATED ANTIBIOTICS ON THE IMMUNE RESPONSE

Preclinical studies evaluating microencapsulated antibiotics in animals have demonstrated that targeted local release of antibiotics directly into infected soft tissue and bone via sustained release of the drug from poly DL(lactide-co-glycolide) will greatly enhance antibiotic efficacy for both prophylaxis and treatment. Antibiotic hypersensitivity was, from the beginning, the most obvious untoward clinical concern of this novel approach to antibiotic delivery. What effect would sustained antibiotic release have on the hypersensitive patient?

Prior to the filing of applicants' parent application Serial No. 590,308 on March 16, 1984, which disclosed the local application of encapsulated antibiotics to treat wound infection, it was commonly known that an inherent property of free antibiotics such as ampicillin, is that they elicit an immune response in man and induce the production of antibodies. Thus, interest in the immune response elicited from the sustained release of immunogens intensified in order to capture the beneficial aspects of this event immunogenic event in a manner which would advance the frontiers of medical science. This led to additional studies with sustain released antibiotics and led the inventors to postulate that antigens encapsulated in lactide/glycolide could potentially

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provide a more effective method of active immunization than free antigen
alone. In follow on experiments, vaccine antigens were encapsulated and
alone. In follow on experiments, the studies were performed to explore this hypothesis as illustrated in Phase II,
herein (Phase II).

VI. BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the effect of microencapsulated ampicillin (MEAA) on the immune response when mice are treated with free ampicillin, ampicillin encapsulated within biodegradable-biocompatible microspheres and placebo poly (Lactide/glycolide) microspheres, by measuring the specific IgG antibodies to free ampicillin and MEAA in sera of treated mice by ELISA.

Figure 2 shows that guinea pigs sensitized with free or microencapsulated ampicillin developed specific IgG antibodies to ampicillin as measured by ELISA.

Figure 3 shows the in vitro release of [14C]-ampicillin anhydrate from sterilized microcapsules/spheres (45 to 106 micrometers in diameter) into 0.1 molar potassium phosphate 2 receiving fluid (pH 7.4) maintained at 37°C. The microcapsules 3 consisted of about 10 weight percent ampicillin anhydrate and 4 about 65 weight percent 53:47 DL-PLG polymer. 5 6

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- Figure 4 shows the in vitro release of [14C]-ampicillin anhydrate from sterilized microcapsules (10 to 100 micrometers 7 consisting of about 35 weight percent ampicillin and about 65 8 weight percent of 53:47 DL-PLG polymer. 10
 - Figure 5 shows the mean daily excretion of [14C] from sterilized injections of 11 subcutaneous microencapsulated and unencapsulated [14C]-ampicillin anhydrate. 12 13
 - Figure 6 illustrates that encapsulated as well as the ampicillin anhydrate showed a fast release of drug during Day 1. 14 By Day 4, the amount of ampicillin found in the serum of animals 15 dosed with the unencapsulated drug was below the level of 16 detection of the assay, whereas serum levels of ampicillin were 17 dectable in animals receiving encapsulated ampicillin for mp to 11 18 19 days. 20
 - Figure 7 shows mean serum levels of ampicillin at 1-hour following implantation of either microencapsulated ampicillin or 21 22

- unencapsulated ampicillin into the medullary canal of the rabbit
- tibia with experimental osteomyelitis.
- Figure 8 shows the mean serum Serum Cefazolin Levels. concentrations of cefazolin that were measured at 1 hour and 24 3 hours following local antibiotic therapy with either CZ 4 microspheres (Group A) or free CZ powder (Group B) in the rabbit 5 fracture-fixation model. At 1 hour, the mean serum cefazolin levels were approximately 32 times higher for the Group B animals 7 who had received local antibiotic therapy with free CZ powder (18.7 8 ± 6.1 ug/ml) as compared to the Group A animals who were treated 9 with CZ microspheres)0.57 \pm 0.27 ug/ml). This difference in the 10 mean serum cefazolin levels between the two groups was 11 statistically significant (p = 0.0023) by Student's t test. At 24 12 hours following local treatment, no cefazolin was detected in the 13 sera of the rabbits who had received free CZ powder (Group B), 14 however, low cefazolin concentrations were detected in the sera of 15 Group A animals who were treated with the CZ microspheres. It is 16 evident from the data that the free antibiotic diffuses rapidly 17 from the wound and is absorbed into the systemic circulation, 18 whereas, the microspheres remain localized and continue to release 19 low but measurable levels of antibiotic for an extended time 20 21 interval. 22

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Figure 9 shows the size destribution of microspheres wherein the particle size distibution (%) is (a) By number 1-5 (91) and 6-10 (9) and (b) By weight 1-5 (28) and 6-10 (72).

Figure 10shows a scanning electron micrograph of crospheres.

Figures 1(a) and (b) show the In vitro immunization of spleen cells and demonstrates that AF/RI pilus protein remains immunogenic to rabbit spleen cells immunized in vitro after microencapsulation. AF/RI pilus protein has been found to be immunogenic for rabbit spleen mononuclear cells in vitro producing a primary IgM antibody response specific to AF/RI. Immunization with antigen encapsulated in biodegradable, biocompatible microspheres consisting of lactide/glycolide copolymers has been shown to endow substantially enhanced immunity over immunization with the free antigen. To determine if microencapsulated AF/RI maintains the immunogenicity of the free pilus protein, a primary in vitro immunization assay was conducted. Rabbit spleen mononuclear cells at a concentration of 3x10° cells/well.

Triplicate wells of cells were immunized with free AF/RI in a dose range from 15 to 150 ng/ml or with equivalent doses of AF/RI contained in microspheres. Supernatants were harvested on days 7, 9, 12, and 14 of culture and were

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Supernatant control values were subtracted from those of the immunized cells.

Cells immunized with free pilus protein showed a significant positive IgM response on all four days of harvest, with the antibody response increasing on day 9, decreasing on day 12, and increasing again on day 14. Cells immunized with microencapsulated pilus protein showed a comparable positive IgM antibody response as cells immunized with free pilus protein. In conclusion, AF/RI maintains immunogenicity to rabbit spleen cells immunized in vitro after microencapsulation.

Figures 1 2a) and (b) show in vitro immunization of Peyer's patch cells. Here the AF/RI pilus protein remains immunogenic to rabbit Peyer's patch cells immunized in vitro after microencapsulation. AF/RI pilus

patch cells. Here the AF/RI pilus protein remains immunogenic to rabbit Peyer's patch cells immunized in vitro after microencapsulation. AF/RI pilus protein has been found to be immunogenic for rabbit Peyer's patch mononuclear cells in vitro producing a primary IgM antibody response specific to AF/RI. Immunization with antigen encapsulated in biodegradable, biocompatible microspheres consisting of lactide/glycolide copolymers has been shown to endow substantially enhanced immunity over immunization with the free antigen. To determine if microencapsulated AF/RI maintains the immunogencity of the free pilus protein, a primary in vitro immunization assay was conducted. Rabbit Peyer's patch mononuclear cells at a concentration of 3x10° cells/ml were cultured in 96-well, round bottom microculture plates at a final concentration of 6x10° cells/well. Triplicate wells of cells were immunized with free AF/RI in a dose range from 15 to 150 ng/ml or with equivalent dose of AF/RI contained in microspheres. Supernatants were

harvested on days 7, 9, 12, and 14 of culture and were assayed for free

AF/RI pilus protein specific IgM antibody by the ELISA. Supernatant control

values were subtracted from those of the immunized cells. Cells immunized

with free pilus protein showed a significant positive IgM response on all four

days of harvest, with the highest antibody response on day 12 with the highest

antigen dose. Cells immunized with encapsulated pilus protein showed a

positive response on day 12 with all three antigen doses. In conclusion, AF/RI

pilus protein maintains immunogenicity to rabbit Peyer's patch cells

immunized in viito after microencapsulation.

Figure 13hows proliferative responses to AF/RI by rabbit Peyer's patch cells. Naive rabbits were primed twice with 50 micrograms of either non-encapsulated (rabbits 132 and 133) or microencapsulated (rabbits 134 and 135) AF/RI pili by endoscopic intraduodenal inoculation seven days apart. Seven days following the second priming, Peyer's patch cells were cultured with AF/RI in 96-well plates for four days followed by a terminal six hour pulse with [3H]thymidine. Data shown is the SI calculated from the mean cpm of quadruplicate cultures. Responses were significant for all rabbits: 132 (p=0.013), 133 (p=.0006), 134 (p=0.0016), and 135 (p=0.0026). Responses were significantly different between the two groups. Comparison of the best responder in the nonencapsulated antigen group (rabbit 133) with the lowest responder in the microencapsulated antigen group (rabbit 134) demonstrated an enhanced response when the immunizing antigen was microencapsulated (p=0.0034).

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Additionally, Figure 1 3relates to the in vitro lymphocyte
of rabbit lymphoid tissues with encapsulated th
proliferation after sensitization of F. coli strain RDEC-1. The AF/RI non-encapsulated AF/RI pilus adhesion of E. coli strain RDEC-1. The AF/RI
non-encapsulated AF/RI pilus adnesson of annual that allows RDEC-1 to
non-encapsulated AF/RI phus and an adherence factor is a plasmid encoded pilus protein that allows RDEC-1 to
haush borders. We investigated and
of encapsulating purified A172
non-reactive microspheres composed of polymerized lactide and glycolide,
mon-reactive microspheres composed of post- materials used in resorbable sutures. The microspheres had a size range of
materials used in resorbable sutures. The interespitation
relected for Peyer's Patch localization, and
NZW rabbits were immunized two
0.62% protein by weight. N24 to the open content of the protein by weight. N24 to the open content of the protein by weight. N24 to the open content of the protein by weight. N24 to the open content of the
micrograms of either encapsulated or non-encapsulated or non-encap
AF/RI by intraduodenal inoculation
visconian respone to purified ARVRI was consessed in
Lymphocyte profiteration in temporal at seven days and showed that encapsulating the antigen into microspheres
at seven days and showed that encapsular at seven days are seven days and showed that encapsular at seven days are seven days and showed that encapsular at seven days are seven days and showed that encapsular at seven days are seven days and showed that encapsular at seven days are seven days are seven days at the seven days are s
at seven days and snowed that of the Peyer's Patch; however, no enhanced the cellular immune response in the Peyer's Patch; however, no
as observed in spleen or mesenteric sympathic
significant increase was observed and significant increase was observe
data suggest that encapsulation of Astronomy
immune response.
d show proliferative responses to Al 122 symmetry
Peptides by rabbit Peyer's patch cells. Naive rabbits were primed twice with
peptides by rabbit Feyer a partial period (rabbits 132 and 133) or
peptides by rabbit Feyor of 50 micrograms of either non-encapsulated (rabbits 132 and 133) or
50 micrograms of class to the second microencapsulated (rabbits 134 and 135) AF/RI pili by endoscopic microencapsulated (rabbits 134 and 135) AF/RI pili by endoscopic
enten days apart. Seven days some training
intraduodenal inoculation seven deposition intraducent deposition in the intraducent deposition deposition in the intraducent deposition depo
priming, Peyer's Pauli Control

. 5

	(Fig142), AF/R1 79-94 (Fig.16), AF/R1 108-123 (Fig.4c), or AF/R1
	(Fig14a), AF/R1 79-94 (Fig.16d), reserved for four days followed by a terminal 40-47/79-86 (Fig.16d) in 96-well plates for four days followed by a terminal
	40-47/79-86 (Fig.16d) in 96-well plates for four the SI calculated from the
	40-47/79-86 (Fig. 61) in 90 to the six hour pulse with [3H]thymidine. Data shown is the SI calculated from the six hour pulse with [3H]thymidine.
	mean cpm of quadruplicate cultures. The responses of rabbits 132 and 133
5	mean cpm of quadruplicate cultures. Were not significant to any of the peptides tested. Rabbit 134 had a significant were not significant to any of the peptides tested. Rabbit 134 had a significant
6	2001 40.55 (n=0.0001), (b) AF/RI 19-74 (P
	sarro 86 (n=0.025), but not to (c) AP/RI 100-125.
7	is and remonse to (a) AF/R1 40-55 (p=0.034), (b) AF/R1
8	AE/R1 108-123 (p < 0.0001), but not to (d) /12/11
9	m's demonstrates enhanced proliferative response to per-
10	and its mucosal priming with microencapsulated pint.
11	PDEC. 1 attachment to rabbit intestinal brush dollars.
12	were selected by theoretical criteria from the
13	amino acid peptides were served. AF/RI 40-55 sequence as probable T or B cell epitopes and were synthesized: AF/RI 40-55
14	as a B cell epitope, 79-94 as a T cell epitope, and 108-123 as a T and B cell
15	as a B cell epitope, 79-94 as a 1 cell epitope. We used these peptides to investigate a possible immunopotentiating epitope. We used these peptides to investigate a possible hipcompatible
16	epitope. We used these pepudes to investigate a september and the pepudes and the pepudes are september and the pepudes and the pepudes are september and the pepudes and the pepudes are september and the pepude
17	epitope. We used these populations and short of encapsulating purified Af/RI pili into biodegradable, biocompatible effect of encapsulating purified Af/RI pili into biodegradable, biocompatible
18	effect of encapsulating partitions of polymerized lactide and glycolide at a size range microspheres composed of polymerized lactide and glycolide at a size range microspheres composed of polymerized lactide and glycolide at a size range
19	that promotes localization in the Peyer's Patch (5-10 micrometers). NZW
20	rabbits were primed twice with 50 micrograms AF/RI by endoscopic
	incomprise and their Peyer's Patch Eeus well of the Peyer's Patch
21	In two rabbits which had received the property of the property
22	ambiferation was observed to AFIRI 40-33
23	AF/RI, lymphocyte promotes both rabbits and to 108-123 in one of two rabbits. No responses to any of the
24	DOM Impage Tra

	peptides were observed in rabbits which received non-encapsulated AF/RI.
1	peptides were observed in rabbits which the representation of AF/RI may enhance the cellular These data suggest that encapsulation of AF/RI may enhance the cellular
2	
3	response to peptide antigens. Figures 1 5a-d show B-cell responses of Peyer's patch cells to
4	Figures 1 5 -d show B-cell responses 6.5 5.7
5	AF/R1 and peptides.
6	AF/R1 and peptides. Figures 162-d show B-cell responses of Peyer's Patch cells to
7	AF/R1 and peptides.
8	AF/R1 and peptides. Figures 1 2-d show B-cell responses of spleen cells to AF/R1 and
9	Peptides.
10	Peptides. Figures 18 a-d show B cell responses of spleen cells to AF/R1
11	and peptides.
12	and peptides. Figures 15 through 18, illustrate enhanced lymphocyte antibody
13	response by mucosal immunization of rabbits with microencapsulated AF/R1
	The AF/RI pilus protein has been found to be immost be
14	a power's patch cells in vitro producing a primary again
15	The purpose of this study was to determine the
16	is enhanced by microencapsulation.
17	hiodegradable, biocompatible microspheres competed
18	tide approximents, had a size range of 3-10 interometer.
19	lactide-glycolide copolymetry containing 0.62% pilus protein by weight. Initially, NZW rabbits were
20	immunized twice with 50 micrograms of either encapsulated or
21	immunized twice with 50 interegrations immunized twice with 50 interegrations and immunized twice with 50 interegrations are immunized twice with 50 interegrations and immunized twice with 50 interegrations are immunized twice with 50 interegrations and immunized twice with 50 interegrations are immunized with 50 interegrations are imm
22	non-encapsulated AF/RI via intraduced and in microculture at final volume challenge, 6x10 ^s rabbit lymphocytes, were set in microculture at final volume
23	challenge, 6x10 ^s rabbit lympnocytes, was a series of three different synthesic 16
24	of 0.2 ml. Cells were challenged with AR/RI or three different synthesic 16

	is a sendicted T. B or T and B cell
	amino acid peptides representing, either predicted T, B or T and B cell
2	epitopes in a dose range of 15 to 150 ng/ml for splenic cells or 0.05 to 5.0
3 .	piccomms/ml for Peyer's patch mononuclear cells (in triplicate).
	Supernatants were collected on culture days 3, 5, 7, and 9 assayed by ELISA
4	for anti-AF/R1 antibody response as compared to cell supernatant control.
5	Significant antibody responses were seen only from spleen and Peyer's patch
6	Significant antibody responses were personal significant antibody cells from rabbits immunized with microencapsulated AF/R1. The antibody
7	cells from rabbits immunized with microchapsurers mainly an IgM response.
8	response tended to peak between days 5 and 9 was mainly an IgM response.
9	The results for the predicted epitopes were similar to those obtained with
	purified AF/RI. In conclusion, intestinal immunization with AF/RI pilus
10	protein contained within microspheres greatly enhances both the spleen and
11	Peyer's patch B-cell responses to predicted T & B-cell epitopes.
12	Peyer's patch B-cell responses to AF/R1 40-55 by Figure 19, shows proliferative responses to AF/R1 40-55 by
13	Figure 19, snows prometable 19.
14	rabbit MLN cells. Naive rabbits were primed twice with 50 micrograms of
15	either nonencapsulated (rabbits 132 and 133) or microencapsulated (rabbits 134
	and 135) AF/R1 pili by endoscopic intraduodenal inoculation seven days apart.
16	Seven days following the second priming, MLN cells were cultured with
17	AF/R1 40-55 for four days in 24-well plates. Cultures were transferred into
18	AF/R1 40-55 for four days in Days in Days in Physician pulse. Data shown is the SI 96-well plates for a terminal [3H]thymidine pulse. Data shown is the SI
19	96-well plates for a terminal [Prijulyment purples Responses of rabbits
20	calculated from the mean cpm of quadruplicate cultures. Responses of rabbits
21	132 and 133 were not statistically significant. Responses were significant for
	rabbits 134 (p=0.0.0051) and 135 (p=0.0055).
22	Figure 20 shows proliferative responses to AF/R1 40-53 by
23	rabbit spleen cells. Naive rabbits were primed twice with 50 micrograms of
24	rabbit spleen cens.

•
either nonencapsulated (rabbits 132 and 133) or microencapsulated (rabbits 134
and 135) AF/R1 pili by endoscopic intraduodenal inoculation seven days apart.
and 135) AF/R1 pili by endoscopic intranscentistic cells were cultured with
and 135) AF/R1 pill by Elitoscopic and 135) AF/R1 p
as the four days in 24-well plates. Cultures with
in this wind pulse. Data site of the same
of quadruplicate cultures.
calculated from the mean control of queen calculated from the mean calculated from the mean calculated from the property of queen calculated from the mean calculated from the property of queen calculated from the queen calculate
132 and 133 were not stansucary as (=0.0066)
rabbits 134 (p=0.0.0005) and 135 (p=0.0066).
rabbits 134 (p=0.0.0005) Electrical Figure 24. A. SDS-PAGE of intact CFA/I (lane 1), trypsin
treated CFA/I (lane 2), and S. aureus V8 protease treated CFA/I. Molecular
ciadioidual hands were estimated from molecular weight
(on left). Multiple lanes of both trypsin and V8 treated CFA/I were
(on left). Multiple lates of the transferred to PVDF membranes where bands corresponding to the
transferred to PVDF membranes where the transferred to the transferred t
approximate molecular masses of 3500 (trypsin digest, see arrow lane 2) and
approximate molecular interests and subjected to Edman 6000 (V8 digest, see arrow lane 3) were excised and subjected to Edman 6000 (V8 digest, see arrow lane 3)
24 Parent sing sequence of protein fragments from date
degradation. B. Resulting Experiment in the intact protein. Underlined, (position of sequenced portion of fragment in the intact protein.
(position of sequence property
italisized residues are amino acted and the continuous sera of

Figure 25. ELISA assay results testing hyperimmune sera of monkeys (A)2Z2 (monkey 3), (B) 184(D) (monkey 1) and (C) 34 (monkey 2) to CFA/I primary structure immobilized on polyethylene pins. Monkey sera diluted 1:1000. Peptide number refers first amino acid in sequence of octapeptide on pin from CFA/I primary structure OD 405 refers to optical density wavelength at which ELISA plates were reat (405 nm).

. 16

Figure '26 Complete sequence of CFA/I (147 amno acids) with
Deall excernition site (boxed areas) as defined by each individual monkey
Derived from data in Figure 23.
Figures 27-29 Lymphocyte proliferation to symmetry
of CEAT. Each monkey was immunized with three i.m.
CEAT subunits in adjuvant, and its spleen cells were consider
is anotheric decapeptides which had been constructed using the Pepsear
The decapeptides represented the entire CFA/I protein.
siens of synthetic peptide used included 6.0, 0.6, and 0.00
Values shown represent the maximum profiterative response
the env of the three concentrations of antigen used + the standard
The come of the control peptide for each of the three monkeys was
$\frac{1}{100}$ + $\frac{1}{100}$ and $\frac{1}{100}$
$1,518 \pm 30,951 \pm 20,000$ control for each of the three monkeys was $1,319 \pm 60,325 \pm 13$, and $1,951$
o . a
± 245 respectively. Figures 30-32 Lymphocyte proliferation to 6.0, 0.6, and 0.06
contest conthetic decapeptides of CFA/I in one monkey. The monkey
micrograms/mi synthetic descriptions of CFA/I subunits in adjuvant, (222) as immunized with three i.m. injections of CFA/I subunits in adjuvant,
and its spleen cells were cultured with synthetic decapeptides which had been
and its spleen cells were curious. The decapeptides represented the constructed using the Pepscan technique. The decapeptides represented the
entire CFA/I protein. Values shown represent the proliferative response which
entire CFA/I protein. Values 2.000 p
occurred to 6.0 micrograms/ml (Fig. 32) of antigen ± the standard deviation. The cpm of
0.06 micrograms/ml (Fig. 32) of The

	control Was
1	the control peptide was 1,553 ± 33 and the cpm of the media control was
2 .	1,951 ± 245. Figure 33 shows that rabbits numbers 21 and 22 received
3	Figure 33 shows that resolves at doses of AF/R1 of 200 intraduodual administration of AF/R1 microspheres at doses of AF/R1 of 200
4	intraduodual administration of AF/RI masterprise and 21 then sacrificed on micrograms (ug) on day 0 and 100 ug on day 7, 14, and 21 then sacrificed on
5	micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 0 and 100 ug ct. — 3 micrograms (ug) on day 10 in day 31. The spleen, Peyer's patch and ileal lamina propria cells at 6 x 10 in day 31. The spleen, Peyer's patch and ileal lamina propria cells at 6 x 10 in day 31.
6	day 31. The spleen, Peyer's patch and MEM and AF/R1 1-13, 40-55, 0.2 ml in quadriplate were challenged with AF/R1 and AF/R1 1-13, 40-55,
7	0.2 ml in quadriplate were challenged with 12.0 ml and 15 ug/ml 79-94, 108-123, and 40-47, 79-85 synthetic peptides at 15, 1.5 and .15 ug/ml
8	79-94, 108-123, and 40-47, 79-85 synthesis popular for 4 days. The supernatants were tested for IL-4 using the IL-4/IL-2
9	for 4 days. The supernatants were used for 4 days.
10	dependent cell line cells CT4R at 50,000, were supernatant for 3 days then pulsed with tritiated thymidine for 4 hrs, cells
11	supernatant for 3 days then pulsed with a research and harvested and the tritiated thymidine incorporation determined, averaged and harvested and the tritiated thymidine incorporation determined, averaged and
12	harvested and the tritiated thymidine interpretable harvested and the tritiated for the property of the property o
13	expressed with one standard deviation the standard deviation (log CFU/gm) in Figure 34 shows that RDEC-1 colonization (log CFU/gm) in
14	Figure 34 shows that RD20 1 000.
15	cecal fluids was similar in both groups (mean 6.3 vs 7.3; p=.09). Figure 35 shows that rabbits given AF/R1-MS remained well
16	Figure 35 shows that rabbits grown and 4/6 gained weight after challenge, whereas 9/9 unvaccinated rabbits lost
17	and 4/6 gained weight after challenge, whereas and 4/6 gained weight after challenge (mean weight change +10 vs -270 grams p<.001).
18	weight after challenge (mean weight charge very weight after challenge (mean weight charge very the second of RDEC-1 attachment to
19	Figure 36 shows that the interest of CFA/II microsphere
20	the cecal epithelium was zero in Vacculates, Lie and the cecal epithelium was zero in Vacculates, Lie and the cecal epithelium was zero in Vacculates, Lie and the cecal epithelium was zero in Vacculates, Lie and Li
21	Figure 37. Particle size distribution of number or volume verses
22	vaccine Lot L74F2 values are percent frequency of number or volume verses
23	distribution. Particle size (diameter) in microns. 63% by volume are between
24	5-10 um and 88% by volume are less then 10 um.

	Figure 38. Scanning electron photomicrograph of CFA/II
1	Figure 38. Scanning steed on a supersents 5 um distance.
2	microsphere vaccine Lot L7472 standard bar represents 5 um distance.
3	Figure 39. Twenty-two hour CFA/II release study of CFA/II
4	microsphere vaccine Lot L7472. Percent cumulative release of CFA/II from
5	three sample: A, 33.12 mgm; B, 29.50 mgm , 24.20 mgm at 1, 3, 6, 8, 12
	and 22 hour intervals. Average represents the mean ± ISD.
6	Figure 40. Serum IgG antibody reponse to CFA/II microsphere
7	vaccine Lot L7472 following 2 25 ug protein IM immunization on
8	day 0 in 2 rabbits. Antibody determines on serial dilution of sera by ELISA
9	day 0 in 2 rabbits. Antibody determines on the
10	and expressed as mean titer versus day 0, 7 and 14.
11	Figure 41. Serum IgG antibody response to CFA/II
12	microsphere vaccine Lot L7F2 following 2 25 ug protein IM
13	immunizations on day 0 if rabbit 107 & 109. Antibody determined on serial
	dilution (in duplicate) of sera by ELISA and expressed as mean titer versus day
14	
15	0, 7 and 14. Figure 42. Lymphocyte proliferative responses for Peyer's
16	patch cells of rabbits 65 (figure 47 (a)), 66 (figure 42 (b)), 83 (figure 42 (c)),
17	patch cells of rabbits 63 (figure 47 (a)), 65
18	86 (figure 42 (d)), and 87 (figure 42. (e)) immunized intraduodenally with 50
19	mgm protein of CFA/II microsphere vaccine 4 and 7 days earlier. The cells
20	are challenged in vitro with CFA/II or BSA at 500, 50 and 5 ug/ml or media
	is emplicate. The uptake of tritiated thymidine in Kcp is expressed as mean +
21	Ten. Heing the paired student t-test, the p values of 500 ug/ml dose of
22	CFA/II compared to media control are: 65,p = 0.0002; 66,p = 0.0002; 83,p
23	
24	= 0.0002; and 86, $p = 0.0002$.

	Figure 43 Lymphocyte proliferative responses from Peyer 3
1	Figure 43 Lymphocya P (5 sure 43 (b)), 80 (figure 43 (c)),
• 2 .	Figure 43 Lymphocyte P patch cells of rabbits 77 (figure 43(a)), 78 (figure 43 (b)), 80 (figure 43 (c)), patch cells of rabbits 77 (figure 43(a)), 78 (figure 43 (b)), 80 (figure 43 (c)),
	(5 cm (3 (e)) 1mmun25
3	A Land VACCING 14 MILE
4	or a with CFA/H of Doil
5	the of the plate.
6	15D Using the party
7	CEA/II compared to
· 8	protein of 500 ug/ml dose of C178889 protein of 500 ug/ml dose of
9	0.0001; 78; = 0.0015; 80, p
10	= 0.0001. Figure 44. ELISPOT assay of spleen cells from rabbits 65
11	Figure 44. ELISPOT 23.45 (c)), 86 (figure 44 (d)), and
12	Figure 44. ELISTOT LT. (figure 44 (a)), 66 (figure 44. (b)), 83 (figure 44 (c)), 86 (figure 44 (d)), and
13	(figure 44 (a)), 66 (figure 44 (e)) immunized intraduodenally with 50 mgm protein 87 (figure 44 (e)) immunized intraduodenally with 50 mgm protein These were cells placed
	ing 14 and 7 days carner.
14	$\frac{1}{2}$ and $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$ $\frac{1}{2}$
15	anihodies specific for CFA/II anugen.
16	less cells versus culture day to
17	TI ICOUT ASSAU OI SPICE!
18	The colls were placed into microculture and matter
19	A B by FI ISPO1 for cent according
20	on days 0, 1, 2, 3, 4 and 5 by 22225 of the for CFA/II antigen. The results are expressed as number per 9 x 10' sphere.
21	for CFA/II antigen. The results are expenses
22	cells versus culture day tested.
	rigure 46. Curve for determining vaccination dosages for
23	regimen b.
24	ickimer .

1	Figure 47 Hepatitis B surface antigen release from 50:50 poly
	(DL-lactide-co-glycolide).
3	Figures 19 and 20 serve to illustrate that inclusion of
4	Escherichia coli pilus antigen in microspheres enhances cellular
5	immunogenicity.
	wolease from a
6	FIG.48 shows a comparison of drug release from a
7	worsys a controlled release 5,500
8	tional administrations
9	valley levels from conventional
10	
11	release administration. FIG. 49 shows a scanning electron micrograph of PLGA FIG. 49 shows a scanning electron micrograph of PLGA
12	FIG. 49 shows a scanning determined in the invention microspheres prepared by the process described in the invention
13	using 50/50 uncapped polymer of Mw 8-12k dalton and shows
14	using 50/50 uncapped polymer of superior sphere morphology, sphere integrity, and narrow size
15	distribution.
16	FIG.49 a shows a scanning electron micrograph of PLGA microspheres prepared by conventional solvent evaporation method microspheres prepared by conventional solvent evaporation.
1.7	microspheres prepared by conventioned Mw 8-12k dalton.
18	using a 50/50 uncapped polymer of Mw 8-12k dalton.
19	rig.50 shows cumulative Histatin release from PLGA
20	microspheres, wherein release profiles from several batches are
21	and and anned bolymer (
22	narameters are valled
23	between 1 and 100 days. FIG. 51 shows a scanning electron micrograph of solid, smooth
24	FIG. 51 shows a scanning electron mid-19-1-

- spherical surfaces of PLGA microspheres prepared by the method of
- in the invention using 50/50, end-capped polymer (of Mw 30-40k
- dalton). 3
- FIG. 52 shows cumulative Histatin release from PLGA
- microspheres, wherein the release profiles are from several 4 5
- batches prepared using 50/50, uncapped and end-capped polymer of
- Mw 30-40k daltons, and wherein the process parameters are varied
- to modulate release between 28 to 60 days. 8
- FIG. 53 shows cumulative Histatin release from PLGA 9
- microspheres, wherein combined release profiles from several 10
- batches have been prepared using 50/50, uncapped and end-capped
- 12 polymer of Mw 8-40k daltons, while varying the process parameters 11
- to modulate release between 1 and 60 days.
- FIG. 54 shows a cumulative percent release of LHRH from PLGA
- 15 microspheres prepared using uncapped polymer of Mw 8-12 daltons.

	VII. DETAILED DESCRIPTION OF THE INVENTION
1	VII. DETAILED DESCRIPTION of active core materials, This invention relates to the encapsulation of active core materials,
2	This invention relates to the encapsulation to the mammalian animal
3	especially those which are medically beneficial to the mammalian animal
4	kingdom, such as biologically active agent(s), drug(s), or substance(s) within a
5	biodegradable-biocompatible polymeric matrix.
6	More precisely, applicants have discovered a medicinally beneficial
7	tit and process with the following itemized features:
8	for the burst-free, sustained, programmable reference
	1/2) ever a period from 1-100 days, which comprises. (1)
9	A carrier which may contain pharmaceutically acceptance
10	adjuvant, comprised of a blend of uncapped and end-capped biodegradable-
11	
12	biocompatible copolymer. 2. The composition of Item 1 wherein the polymeric substance is
13	
14	poly(lactide/glycolide).
15	poly(lactide/glycolide). 3. The composition of Item 2, wherein the poly(lactide/glycolide) is a blend i
16	of uncapped and end-capped forms, in ratios ranging from 100/0 to 1/99.
17	4. The composition of Item 3 wherein the copolymer (lactide to glycolide
18	and end-capped polymer is 90/10 to 40/00.
19	The composition of Item 4 wherein the copolymer (lactide to give lactide to gi
	and end-capped polymer is 46/32 to 32 ver
20	L/G) ratio for uncapped and the second second ratio of the copolymer. 6. The composition of Item 2 wherein the molecular weight of the copolymer.
21	a ann an ann an an an an an an an an an
22	7. The composition of Item 3 wherein the active material is biologically active
23	7. The composition of near 3 where 3
24	agent.
25	agent. 8. The composition of Item 7 wherein the agent is selected from the group
26	consisting essentially of antibacterial agents; peptides; polypeptides;
27	antibacterial peptides; antimycobacterial agents; antimycotic agents; antiviral

	-40°
1	agents; hormonal peptides; cardiovascular agents; narcotic antagonists;
2	steroids including HIV dictated and
	inhibitors) and AZT; estrogens; progesums, general
3	is contest non-steroidal anti-inflammatory agents, parasympton
4	tranquilizers; decongestants, decongestants,
5	because non-extrogenic and non-progestional steroids, sympassion
6	nutrients; anti-migraine drugs, electrony
7	replacements; ergot alkaloids; anti-inflammary agents; prostaglandins;
8	antibodies; enzymes; growth factors,
9	heromones; prodrugs; psychotropic diugs, incomis
10	a wine drives: ametite suppressants/stimulants and company
11	antiblood clotting trugs, appearance thereof; contraceptive agents include estrogens such as diethyl silbestrol; 17-
12	beta-estradiol; estrone; ethinyl estradiol; mestranol; progestins such as
13	norethindrone; norgestryl; ethynodiol diacetate; lynestrenol;
14	medroxyprogesterone acetate; dimethisterone; megestrol acetate;
15	medroxyprogesterone acetate; norgestimate; norethisterone; ethisterone; melentate; chlormadinone acetate; norgestimate; norethisterone; ethisterone; and
16	norgestimate; norethisterone; ethisterone; melengestrol; norethynodrel; and
17	norgestimate; norethisterone, such as nonyphenoxypolyoxyethylene glycol; spermicidal compounds such as nonyphenoxypolyoxyethylene glycol;
18	spermicidal compounds such as non-
19	such as aluminum hydroxide; calcium carbonate; magnesium carbonate;
20	such as aluminum hydroxide; careful enterest and antifertility agents; sodium carbonate and the like; non-steroidal antifertility agents;
21	sodium carbonate and the like, how server sodium carbonate and the like, how server se
22	parasympathomimetic agents; psychodizary such as chloropromaquine HCL; clozapine; mesoridazine; metiapine;
23	such as chloropromaquine HCL; clozaphio, more such as chlordiazepoxide;
24	reserpine; thioridazine; minor tranquilizers such as chlordiazepoxide;
25	diazepam; meprobamate; temazepam and the like; rhinological decongestants;
26	diazepam; meprobaniate, continued in the diazepam; meprobaniate, sodium pentobarbital; sedative-hypnotics such as codeine; phenobarbital; sodium pentobarbital;
27	sedative-hyphotics seem as testosterone and testosterone sodium secobarbital; other steroids such as testosterone and testosterone

	· vm/ conte
1	nutrient such as the essential amino acids; essential fats; anti-HIV agents;
2	tine A 7T: antimalarials such as 4-aminoquinolines; 8 aminoquinomies,
3	pyrimethamine; anti-migraine agents such as mazindol; pnenterninte, and
4	and such as L-dopa; antispasmodics such as arropure,
	methscopolamine bromide; antispasmodics and anticholingeric agents such a
5	dicertants: enzymes and the like; antitussives such as
6	and noscapine; bronchodilators; cardiovascular agents active
7	as anti-hypertensive compounds; Rauwolfia alkaloids; coronary vasodilators;
8	nitroglycerin; organic nitrites; pentaerythriotetranitrate; electrolyte
9	replacements such as potassium chloride; ergotalkaloids such as ergotamine
10	during the state of the state o
11	methanesulfate; dihydroergocornine methanesulfonate; dihydroergokroyptine
12	methanesulfate; universely methaneusulfate and combinations thereof; alkaloids such as atropine sulfate;
13	methaneusulfate and combanded and combanded analysis; narcotics such as codeine; Belladonna; hyoscine hydrobromide; analgesics; narcotics such as codeine;
14	Belladonna; hyoscine nydrooromae, dihydrocodienone; meperidine; morphine; non-narcotics such as salicylates;
15	dihydrocodienone; meperiume, mospanie dihydrocodienone; meperiume, mospanie antibiotics such as the aspirin; acetaminophen; and d-propoxyphene; antibiotics such as the
16	aspirin; acetaminophen; and d-propoxymenty, cephalosporins including ceflacor and cefuroxime; chloranphenical; gentamicin;
17	cephalosporins including certacor and certacoration; ampicillin; amoxicillin;
18	Kanamycin A. Kanamycin B; the penicillins; ampicillin; amoxicillin;
19	streptomycin A; antimycin A; chloropamtheniol; metromidazole;
20	oxytetracycline penicillin G; the tetracyclines; including minocycline; fluoro-
21	quinolones including ciprofloxacin; ofoxacin; macrolides including
22	quinoloies necessity : clarithromycin; frythromycin; aminioglycosides including gentamicin; clarithromycin; frythromycin; aminioglycosides including ampacillin;
23	amikacin; tobramycin and kanamycin; beta-lactams including ampacillin;
24	polymyxin-B; amphotercin-B; aztrofonam; chloramphenicol; fusidans;
25	polymyxin-B, amphotestary polymyxin-B, ampho
26	systemic antibodies including rifampin; polygenes; sulfunamides; trimethoprin;
27	systemic antibodies including vancomycin; teicoplanin and imidazoles; anti-cancer glycopeptides including vancomycin; teicoplanin and imidazoles; anti-cancer

1	phenobarbital; trimethadione; anti-emetics such as triethylperazine;
2	is incoming such as chlorophinazine; dimenhydrinate; dipnennydianinio,
	becaring: tripelennamine and the like; anti-inflammatory agents such a
3	hammenal agents: hydrocortisone; prednisolone; prednisone; non-normonal
4	agents; allopurinol; for claims water-soluble hormone drugs; antibiotics;
5	antipyretics; analysis,
6	antitumor agents, and antitumor agents, and antitumor agents, anticulcer antitussives; expectorants; sedatives; muscle relaxants; antiepileptics; anticulcer
7	antitussives; expectoration, december, antitussives; expectoration, december, antitussives; antitussives; expectoration, december, antitussives; expectoration, december, antitussives; antitussives; expectoration, december, antitussives; antitussives; antitussives; expectoration, december, decemb
8	agents; antidepressants; antiantergio traggilarits; and antinarcotics; in vasodilators; antihypertensives; diuretics; anticoagulants; and antinarcotics; in
9	vasodilators; antihypertensives, didicator, antihypertensives, didicators; indomethacin; the molecular wight range of 100-100;000 daltons; indomethacin;
10	the molecular wight range of 100-100,000 cares, such as thiotepa; chloramucil; phenylbutazone; prostaglandins; cytotoxic drugs such as thiotepa; chloramucil;
11	phenylbutazone; prostaglandins; cytotoxic diago elemente methotrexate; antigens such as
12	cyclophosphamide; melphala; nitrogen mustard; methotrexate; antigens such as
13	proteins; glycoproteins; synthetic peptides; carbohydrates; synthetic
14	proteins; glycoproteins; synthetic polysaccharides; lipids; glycolipids; lipopolysaccharides(LPS); synthetic
15	polysaccharides, holds, gest a synthetic lipopolysaccharides and with or without attached adjuvants such as synthetic
16	muramyl dipeptide derivatives; antigens of such microorganisms as Neisseria
17	Muschacterium tuberculosis; Picarinii Pnfumonia; riespes vitus
18	2): Hernes zoster; Candidia albicans; Candida dopicans,
19	Trichomonas vaginalis; Haemophilus vaginalis; Group B sureptoccuras cous,
20	hominis: Hemophilus ducreyi; Granuonia inguinato,
	Lamphorathia venerum; Treponema palidum; Brucela aborus Brucela menerum;
21	Campylobacter fetus; Campylobacter fetus;
22	in I estocuira nomona: Listeria monocytogenes; Brucella 0415, Equino
23	herpes virus 1; Equine arteritis virus; IBR-IBP virus; Chlamydia psittaci;
24	herpes virus 1; Equine arents beautiful formation of the second of the s
25	Trichomonas foetus; Taxopiasina generalia abortus eui; Pseudomonas equili; Salmonella abortus ovis. Salmonella abortus eui; Pseudomonas
26	equili; Salmonella abortus ovis. Salmonema aeruginosa; Corynebacterium equi; Corynebacterium pyogenes; Actinobaccilus
27	aeruginosa; Corynebacterium equi, Corynebacterium

Trypanosoma equiperdum; Babesia cabali; Clostridium tetani; antibodies which counteract the above microorganisms; and enzymes including ribonuclease; 1 neuramidinase; trypsin; glycogen phosphorylase; sperm lactic dehydrogenase; 2 sperm hyaluronidase; adenossinetriphosphase; alkaline phosphatase; alkaline 3 phospha esterase; amino peptides; typsin chymotrypsin amylase; muramidase; acrosomal proteinase; diesterase; glutamic acid dehydrogense; succunic and 5 dehydrogenase; beta-glycophosphatase lipase; ATP-ase alpha-peptate gamma-6 glutamyiotranspeptidase; sterold-beta-ol-dehydrogenase; DPN-di-aprorase; and 7 8 combinations thereof. 9. The composition of Item 8 wherein the agent is selected from the group 9 consisting essentially of antibacterial agents; antibacterial peptides; 10 antimycobacterial agents; antimycotic agents; antiviral agents; antiparasitic 11 agents; antifungal; hormonal peptides; cardiovascular agents; narcotic 12 antagonist; analgesics; anesthetics; vaccines; insulins; HIV therapeutic drugs 13 (protease inhibitors); estrogens; progestins; gastrointestinal therapeutic agents; 14 non-steroidal anti-inflammatory agents; parasympathoimetic agents; 15 psychotherapeutic agents; tranquilizers; decongestants; sedative-hypnotics; non-16 estrogenic and non-progestional steroids; sympathomimetic agents; vaccines; 17 vitamins; nutrients; anti-malarial compounds; anti-migraine drugs; electrolyte 18 replacements; ergot alkaloids; analgetics; non-narcotics; anti-cancer agents; 19 anticonvulsants; anti-emetics; antihistamines; anti-inflammary agents; 20 prostaglandins; cytotoxic drugs; antigens; antibodies; enzymes; growth factors; 21 immunomodulators; pheromones; prodrugs; psychotropic drugs; appetite 22 23 suppresants/stimulants; and combinations thereof. 10. The composition of Item 8 wherein the agent is a peptide or polypeptide. 24 11. The composition of Item 10 wherein the agent is a poly peptide. 25 12. The composition of Item 11 wherein the molecular weight of the 26 27

	13. The composition of Item 12 wherein the polypeptide is histatin consisting
1	thering a molecular weight of 1500.
2	of 12 amino acids and having a more of 12 amino acids a more of 12 amino aci
3	14. The composition of Item 1 characteristics of the 1 to 40
4	14. The composition of item 1 calculated and 14. The composition of item 1 to 40 release histatin in an aqueous physiological environment within from 1 to 40 release histatin in an aqueous physiological environment poly(lactide/glycolide)
5	release histatin in an aqueous physical days with a 100/0 blend of uncapped and end-capped poly(lactide/glycolide)
6	days with a 100/0 bleat of 2149. The first than having a L/G ratio of 48/52 to 52/48, and a molecular weight less than
7	
	15,000. 15. The composition of Item 14 wherein the histatin can be completely
8	is in 18 to 40 days and the molecular weight
9	the range of 28,000 w 10,000
10	2 characterized by the carrier
11	16. The composition of Item 2 characters of the histatin in an aqueous physiological environment from 28-70 days
12	90% of the histatin in an aqueous physical poly(lactide/glycolide) having a
13	90% of the histatin in an aqueous physics of the histatin in an aqueou
	x 10 min of 48/52 to 52/48 and a molecular weight range
14	DO 1400 or 151
14 15	
15	daltons.
15 16	daltons. 17. The composition of Item 2 characterized by the capacity to release up to
15 16 17	daltons. 17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days
15 16	daltons. 17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a stempt weight of less than 15,000 daltons.
15 16 17	daltons. 17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons.
15 16 17 18	daltons. 17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons.
15 16 17 18 19	17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons. 18. The composition of Item 13 having analogs of histatin with chain lengths of from 11-24 amino acids of molecular weights from 1,500-3,000 daltons and
15 16 17 18 19 20	17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons. 18. The composition of Item 13 having analogs of histatin with chain lengths of from 11-24 amino acids of molecular weights from 1,500-3,000 daltons and
15 16 17 18 19 20 21 22	17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons. 18. The composition of Item 13 having analogs of histatin with chain lengths of from 11-24 amino acids of molecular weights from 1,500-3,000 daltons and characterized by the following structures:
15 16 17 18 19 20 21 22 23	17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons. 18. The composition of Item 13 having analogs of histatin with chain lengths of from 11-24 amino acids of molecular weights from 1,500-3,000 daltons and characterized by the following structures: 1. D S H A K R H H G Y K R K F H E K H H S H R G Y R
15 16 17 18 19 20 21 22 23 24	daltons. 17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons. 18. The composition of Item 13 having analogs of histatin with chain lengths of from 11-24 amino acids of molecular weights from 1,500-3,000 daltons and characterized by the following structures: 1. D S H A K R H H G Y K R K F H E K H H S H R G Y R 2. K R H H G Y K R K F H E K H H S R
15 16 17 18 19 20 21 22 23	17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons. 18. The composition of Item 13 having analogs of histatin with chain lengths of from 11-24 amino acids of molecular weights from 1,500-3,000 daltons and characterized by the following structures: 1. D S H A K R H H G Y K R K F H E K H H S H R G Y R 2. K R H H G Y K R K F H E K H H S H R G Y R 3. K R H H G Y K R K F H E K H H S R
15 16 17 18 19 20 21 22 23 24	17. The composition of Item 2 characterized by the capacity to release up to 80% of histatin in an aqueous physiological environment from 56-100 days with a 1/99 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 and a molecular weight of less than 15,000 daltons. 18. The composition of Item 13 having analogs of histatin with chain lengths of from 11-24 amino acids of molecular weights from 1,500-3,000 daltons and characterized by the following structures: 1. D S H A K R H H G Y K R K F H E K H H S H R G Y R 2. K R H H G Y K R K F H E K H H S H R G Y R 3. K R H H G Y K R K F H E K H H S R

1	7. KRHHGYKRKF
	*D-amino acid
2	*D-amino acid 19. The composition of Item 10 wherein the biologically active agent is a
3	hermone releasing normans (
4	polypeptide Leutinizing normals respectively. The decapeptide of molecular weight 1182 in its acetate form, and having the
5	
6	STRUCTURE: P-EHWSYGLRPG
7	p-EHWSYGERIO 20. The composition of Item 13 having a molecular weight of from 1,000 to
8	20. The composition of Item 13 having a most
9	250,000 daltons.
10	250,000 daltons. 21. The composition of Item 2 wherein release profiles of variable rates and
11	21. The composition of Item 2 who can be a capped microspheres as a durations are achieved by blending uncapped and capped microspheres as a
12	•
13	cocktail in variable amounts. 22. The composition of Item 2 wherein release of profiles of variable rates and
	22. The composition of Item 2 whereast duration are achieved by blending uncapped and capped polymer in different
14	mherre.
15	a wherein the entrapped polypop
16	torigenic E. Ou (2.23)
17	the vaccine agents against enterousigers group consisting of CFA/I, CFA/II, CS1, CS3, CS6 and CS17, ETEC-related
18	and the second to the second t
19	enterotoxins, and combinations thereof. 24. The composition of Item 23 wherein the entrapped polypeptide consists of
20	24. The composition of Item 23 wherein are of about 800-5000 daltons for
21	24. The composition of Item 25 that 25 that 24. The composition of Item 25 that 25 tha
22	resignic E. coll (E122).
23	of Item 24 wherein the chapped in
	rially of an anugenic system
24	from the group consisting essentially or containing CFA/I pilus protein T-cell epitopes; B-cell epitopes, or mixtures
25	•
26	thereof. 26. The composition of Item 24 wherein the poly(lactide/glycolide) is a bind
27	26. The composition of action

	27. The composition of Item 7 wherein said agent are selected from the group
1	27. The composition of Item 7 where 27. The composition of Item 7
2	consisting of water-soluble hormone trugs, antinustives, expectorants,
3	inflammatory agents, antipyretics, analgesics antitussives, expectorants,
4	antiepileptics, antimiet again,
5	antiarrhythmic drugs, V
6	antiallergic drugs, earthousies, earthousies, earthousies, antinarcotics, in the molecular antihypertensives, diuretics, anticoagulants, antinarcotics, in the molecular
7	maight range of 100-100,000 daltons.
	twherein said biodegranable
8	in an oil phase, and is present in the
9	poly(lactide/glycolide) is in all off prize. 29. The composition of Item 28 wherein concentration of the active agent is in
10	29. The composition
11	the range of 0.1 to about 60% (w/w).
12	the range of 0.1 to about 60% (may) 30. The composition of Item 29 wherein a ratio of the inner aqueous to oil
13	phases is about 1/4 to 1/40(v/v).
14	phases is about 1/4 to 1/40(v/v). 31. The composition of Item 11 wherein the entrapped polypeptide is active at
15	a low pH, such as LHRH, adrenocorticotropic hormone, epidermal growth
16	a climentide is bioactive.
	Size 11 when entrapped polypepado
17	and the stabilizing agent of inolgand design and the stabilizing agent of inolgand design and the stabilizing agent of inolgand design agent des
18	ismin hiological activity of the
19	inner aqueous phase to maintain of the such as 33. The composition of Item 11 wherein when entrapped polypeptide such as 33.
20	33. The composition of Item 11 will be a specific to the composition of Item 11 will b
21	histatin is inactive at a low pH, a non-rouse 60 and Tween 20) and
22	sorbitan fatty acid esters (Tween 80, Tween 60 and Tween 20) and
23	sorbitan fatty acid esters (Tween 60, 2 to 60) polyoxyethylene - polyoxypropylene block copolymers (Pluronics) is added to
24	polyoxyethylene - polyoxypropyrous the inner aqueous phase to maintain biological activity of the released
25	polypeptide.
26	polypeptide. 34. The composition of Item 32 wherein placebo spheres loaded with the H.
27	agents are coadministered with polypepade to a series are coadministered with a series

	activity of the released peptide in instances where the addition of pH-stablizing
1	activity of the released peputic in andesirable for the successful
2	agents in the inner aqueous phase is undesirable for the successful
3	encapsulation of the acid pH sensitive polypeptide.
4	c very 33 wherein placebo spineres very
5	and ministered with polypeputer to and the second s
6	the released peptide where the authors of the released peptide where the
7	surfactants in the inner aqueous phase is undestrable to account
,	supposed by sensitive polypepude.
8	1 comprising a blend of uncapped
9	36. The composition of Item 1 complete solubilization of the copolymer leaves no residual polymer, wherein complete solubilization of the copolymer leaves no residual
10	polymer, wherein complete solutions are concurrently with the complete
11	polymer, wherein complete solution and occurs concurrently with the complete polymer at the site of administration and occurs concurrently with the complete
12	release of the entrapped agent.
13	release of the entrapped agent. 37. A process of using composition of Item 1 for human administration via
14	as intramuscular and subcutations.
15	parenteral routes, such as industries 38. A process of using the composition of Item 1 for human administration
16	via topical route. 39. A process of using the composition of Item 1 for human administration
17	
18	via oral routes. 40. A process of using the composition of Item 1 for human administration
19	40. A process of using the composition routes.
20	via nasal, transdermal, rectal, and vaginal routes. 41. A process of using the composition of Item 1 for human administration in
21	41. A process of using the composition of the respiratory tract.
22	the form of an oral or nasal inhalant for the respiratory tract. the form of an oral or nasal inhalant for the respiratory tract. 42. A process for preparing controlled release compositions characterized by
23	42. A process for preparing controlled research biologically active agents,
24	burst-free, sustained, programmable release of biologically active agents,
25	burst-free, sustained, programmers burst-free, sustained, programmers comprising: Dissolving biodegradable poly(lactide/glycolide), in uncapped comprising: Dissolving biodegradable poly(lactide/glycolide), in uncapped
26	and dissolving a biologically described
27	adding the aqueous layer to the polymer account

	-30°
	w/o emulsion in a solvent-saturated aqueous phase containing a oil-in-water
1	w/o emulsion in a solvent-sate w/o emulsion to an external aqueous layer (o/w) emulsifier; adding said w/o emulsion to an external aqueous layer
2	emuleifier to form a ternary contains
3	containing Oil-in-water contai
4	resulting water-in-oil-in-water (w/o/w) emulsion for sufficient time to remove resulting water-in-oil-in-water (w/o/w) emulsion for sufficient time to remove said solvent, and rinsing hardened microcapsules with water and lyophilizing
5	said solvent, and rinsing hardened hadded
6	said hardened microcapsules. 43. The process of Item 42 wherein a solvent-saturated external aqueous phase
7	43. The process of Item 42 wherein a series to addition of the external
8	is added to emulsify the inner w/o emulsion prior to addition of the external
9	is added to emulsify the liner was a said and the liner was a said added to emulsify the liner was a said and the liner was a said added to emulsify the liner was a said added to emulsion was a said and the liner was a said added to emulsion was a said and the liner was a said added to emulsion was a said and the liner was a said added to emulsion was a said and the liner was a said and
10	between 0.05-500um. 44. The process of Item 42 wherein a low temperature of about 0-4 degree C
11	44. The process of Item 42 wherein a low completion, and a low
12	is provided during preparation of the inner w/o emulsion, and a low
13	is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided during preparation of the temperature of about 4-20 degree C is provided as the temperature of about 4-20 degree C is provided as the temperature of the temperature of about 4-20 degree C is provided as the temperature of the t
14	controlled release compositions
15	burst-free, sustained compositions characterized by burst-free, sustained,
16	- Fhiologically active agents, compression
17	asia polynactide/glycolide) in the
18	dissolving biodegradule poly(about 0) methylene chloride, and dissolving a biologically active agent or active core in
19	water; adding the aqueous layer to the polymer solution and emulsifying to
20	water; adding the aqueous layer to the provide an inner water-in-oil emulsion; stabilizing the w/o emulsion in a provide an inner water-in-oil emulsion; stabilizing the w/o emulsion in a
21	provide an inner water-in-out enturater, solvent-saturated aqueous phase containing a oil-in-water (o/w) emulsifier,
22	adding said w/o emulsion to an external aqueous layer containing oil-in-water
23	adding said w/o emulsion to an external and stirring a resulting water-in-oil- emulsifier to form a ternary emulsion; and stirring a resulting water-in-oil-
24	water (w/o/w) emulsion for sufficient time to remove said solvent; and riming
25	water (w/o/w) emulsion for sufficient and lyophilizing said hardened heardened microcapsules with water; and lyophilizing said hardened
26	
27	microcapsules.

•	46. The process of Item 42 wherein a 100/0 blend of uncapped and end-
1	capped polymer is used to provide release of the active core in a continuous
2	and sustained manner without a lag phase.
3	47. The process of Item 45 wherein a solvent-saturated external aqueous phase
4	is added to emulsify the inner w/o emulsion prior to addition of the external
5	aqueous layer, to provide microcapsules of narrow size distribution range
6	
7	between 0.05-500um.
8	48. The process of Item 45 wherein a low temperature of about 0-4 degree C
9	is provided during preparation of the inner w/o emulsion, and a low
10	temperature of about 4-20 degree C is provided during preparation of the
11	w/o/w emulsion to provide a stable emulsion and high encapsulation efficiency.
12	49. A method for the protection against infection of a mammal by pathogenic
13	organisms comprising administering orally to said mammal an immunogenic
14	amount of an immunostimulating composition consisting essentially of an
15	antigenic synthetic peptide encapsulated within a poly(lactide/galactide) matrix.
16	50. The method of Item 49 wherein the poly(lactide/glycolide) is a blend or
17	uncapped and end-capped forms, in ratios ranging from 100/0 to 1/99.
18	51. The method of Item 49 wherein the poly(lactide/glycolide) is a blend of
19	uncapped and end-capped forms in ratios ranging from 90/10 to 40/60.
	52. The method of Item 49 wherein the infection is a bacterial infection.
20	53. The method of Item 49 where the synthetic peptide contains an epitope
21	selected from the group consisting of CFA/I pilus protein T-cell epitopes, B-
22	cell epitopes or mixtures thereof.
23	54. The method of Item 49 wherein the infection is a viral infection.
24	55. The method of Item 49 wherein the infection is parasitic infection.
25	55. The method of Item 49 wherein the infection is a fungal infection.
26	56. The method of Item 49 wherein the bacterial infection is caused by a
27	57. The method of Item 52 wherein the baseline and second

	-58-
	essentially of Salmonella typhi, Shigella Sonnei, Shigella Flexneri, Shigella
1	estalla boydii Escheria coli, Vibrio choleta,
2	E, Group G, Group I, Group 1, Listeria, Erysipelothrix, Mycobacterium,
3	E, Group G, Group I, Group I, Parties Actinomycetales, Enterobacteriaceae, Vibrio, aeromonas, Aerobic pathogenic Actinomycetales, Enterobacteriaceae, Vibrio, aeromonas,
4	Aerobic pathogenic Actinomycerata, Acineto bacter spo.,
5	Plesiomonas, Helicobacter, W. succinogenes, Acineto bacter spp., Plesiomonas, Helicobacter, W. succinogenes, Acineto bacter spp., Plesiomonas, Helicobacter, W. succinogenes, Acineto bacter spp.,
6	Plesiomonas, Helicobacter, W. Legionella, Brucella, Haemophilus, Bordetalla, Foavobacterium, Pseudomonas, Legionella, Brucella, Haemophilus, Bordetalla,
7	Mycoplasmas, Gardnerella, Streptobacillus, Spirillum, Calymmatobacterium,
8	Mycoplasmas, Gardierena, Ecrelia, Leptospira, Anaerobic Gram-negative Clostridium, Treponema, Borrelia, Leptospira, Anaerobic Gram-negative
9	and Cocci, Anaerobic grant-to-to-
10	Bacilli and Cocci, yersinia, staphylococcus, second
11	Aerococcus, Planococcus, Stormas
	Germella, Pediococcus, Leuconomos.
12	Naisceria Branhamella, Coryne bacterium, campyiooacter, Provincia
13	Placescus Spo., Rhodococcus, Group A. T.
14	the accordance with Item 49 comprising accordance with Item 49 comprising
15	amount of a pharmaceuted
16	to said mammal an immunogenic and to said mammal an immunogenic and the consisting essentially of an antigenic synthetic peptide in the amount of .1 to
17	consisting essentially of all analysis
18	1%. 59. A vaccine for the immunization of a mammal against infection caused by
19	59. A vaccine for the immunization of 2 minor of Item 1.
20	pathogenic organisms prepared from the composition of Item 1.
21	pathogenic organisms prepared 1100 60. The vaccine according to Item 59 wherein the polymeric substance is
22	poly(DL-lactide-co-glycolide).
23	poly(DL-lactide-co-glyconiuc). 61. The vaccine according to Item 60 wherein the relative ratio between the
24	G (G) component is within the range of
	to Item 61 wherein the relative and
25	62. The vaccine according to be amount of lactide and glycolide component is within the range of 90/10 to
26	
27	40/60.

1	63. A vaccine according to Item 62 wherein the pathogenic organisms are
2	bacterial.
3	64. A vaccine according to Item 62 wherein the pathogenic organisms are
4	viral.
5	65. A vaccine according to Item 62 wherein the pathogenic organisms are
6	fungal.
7	66. A vaccine according to Item 62 wherein the pathogenic organisms are
8	parasitic.
9	67. The vaccine according to Item 63 wherein the antigenic synthetic peptide
10	is selected from the group consisting essentially of Synthetic Peptides
11	Containing CFA/I Pilus Protein T-cell Epitopes (Starting Sequence # given)
12	4(Asn-Ile-Thr-Val-Thr-Ala-Ser-Val-Asp-Pro),
13	8(Thr-Ala-Ser-Val-Asp-Pro-Val-Ile-Asp-Leu),
14	12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gin-Ala-Asp),
15	15(Ile-Asp-Leu-Leu-Gln-Ala-Asp-Gly-Asn-Ala),
16	20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
17	26(Pro-Ser-Ala-Val-Lys-Leu-Ala-Tyr-Ser-Pro),
18	72(Leu-Asn-Ser-Thr-Val-Gin-Met-Pro-Ile-Ser),
19	78(Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Gln),
20	87(Gln-Val-Leu-Ser-Thr-Thr-Ala-Lys-Glu-Phe),
21	126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr), and
22	133(Gly-Asn-Tyr-Ser-Gly-Val-Val-Ser-Leu-Val), and
23	mixtures thereof;
24	Synthetic Peptides Containing CFA/I Pilus Protein B-cell (antibody) Eptiopes
25	(Starting Sequence # given)
26	3(Lys-Ana-Ile-Thr-Val-Thr-Ala-Ser-Val),
27	11(Val-Asp-Pro-Val-Idle-Asp-Leu-Leu-Gln-Ala-Asp)

1	32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-Thr-Phe-
2	Glu-Ser-Tyr-Arg-Val),
3	32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe),
4	38(Lys-Thr-Phe-Glu-Ser-Tyr-Arg-Val),
5	66(Pro-Gin-Leu-Thr-Asp-Val-Leu-Asn-Ser),
6	93(Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala),
7	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr),
8	127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
9	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-
10	Ser), and mixtures thereof; and
11	Synthetic Peptides Containing CFA/I Pilus Protein T-cell and B-cell (antibody)
12	Enitones (Starting Sequence # given)
13	3(Lys-Asn-lle-Thr-Val-Thr-Ala-Ser-Bal-Asp-Pro),
14	8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-
15	Ala-Asp),
16	11 (Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
17	20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
18	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-
19	Ser), and
20	126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
21	mixtures thereof.
22	68. The vaccine according to Item 67 wherein the bacteria is selected from the
23	group consisting essentially of Salmonella typhi, Shigella Sonnei, Shigella
24	Flexneri, Shigella dysenteriae, Shigella boydii, Escheria coli, Vibrio cholera,
25	Group D-2, Group E, Group G, Group I, Group 1, Listeria, Erysipelothrix,
26	Mycobacterium, Aerobic pathogenic Actinomycetales, Enterobacteriaceae,
27	Vibrio, aeromonas, Plesiomonas, Helicobacter, W. succinogenes, Acineto

	Haemophilus, Bordetalla, Mycoplasmas, Gardnerella, Streptobacillus,
1	Spirillum, Calymmatobacterium, Clostridium, Treponema, Borrelia,
2	Spirillum, Calymmatovacterium, Leptospira, Anaerobic Gram-negative Bacteria including bacilli and Cocci,
3	Leptospira, Anaerobic Grant-negative Nonsporeforming Bacilli and Cocci, yersinia,
4	Anaerobic gram-Positive Nonspore John Server Aerococcus,
5	staphylococcus, clostridium, Enteroccus, Streptoccus, Aerococcus,
6	Planococcus, Stomatococcus, Micrococcus, Lactoccus, Germella, Pediococcus,
7	Leuconostoc, Bacillus, Neisseria, Branhamella, Coryne bacterium,
8	campylobacter, Arcanobacterium haemolyticum, Rhodococcus spp.
9	Rhodococcus, Group A-4.
•	
10	69. The vaccine according to Item 67 wherein the antigenic synthetic peptide
10	is selected from the group consisting essentially of 4(Asn-Ile-Thr-Val-thr-Ala-
11	
12	Ser-Val-Asp-Pto), 8(Thr-Ala-Ser-Val-Asp-Pto-Val-Ile-Asp-Leu),
13	8(Thr-Ala-Ser-Val-Asp-Leu-Leu-Gln-Ala-Asp), 12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
14	12(Asp-Pro-Val-lie-Asp-Leu-200 Gly-Asn-Ala).
15	15(Ile-Asp-Leu-Leu-Gln-Ala-Asp-Gly-Asn-Ala),
16	20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
17	26(Pro-Ser-ala-Val-Lys-Leu-Ala-Tyr-Ser-Pro),
18	72(Leu-Asn-Ser-Thr-Val-Gin-Met-Pro-Ile-Ser),
19	78(Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Gln),
20	87(Gin-Val-Leu-Ser-Thr-Thr-Ala-Lys-Glu-Phe),
21	126(Ala Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr), and
22	133(Gly-Asn-Tyr-Ser-Gly-Val-Val-Ser-Leu-Val), and mixtures thereof.
22	
	70. The vaccine according to Item 69 wherein the antigenic synthetic peptide is
23	70. The vaccine according to 4(Asn-Ile-Thr-Val-Thr-Ala-ser-Val-Asp-Pro).
24	4(Asn-Ile-Thr-Val-1111-71111-1111-11111-11111-11111-11111-11111-1111

1	71. The vaccine according to Item 69 wherein the antigenic synthetic peptide	:
	is 8(Thr-ala-ser-Val-Asp-Pro-Val-Ile-asp-Leu).	

- 3 72. The vaccine according to Item 69 wherein the antigenic synthetic peptide
- 4 is 12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gin-Ala-Asp).
- 5 73. The vaccine according to Item 69 wherein the antigenic synthetic peptide
- 6 is 15(Ile-Asp-Leu-Leu-Gln-Ala-Asp-Gly-Asn-Ala).
- 7 74. The vaccine according to Item 69 wherein the antigenic synthetic peptide
- 8 is 20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val).
- 9 75. The vaccine according to Item 69 wherein the antigenic synthetic peptide
- is 26(Pro-Ser-Ala-Val-Lys-Leu-Ala-tyr-Ser-Pro).
- 76. The vaccine according to Item 69 wherein the antigenic synthetic peptide
- is 72(Leu-Asn-Ser-Thr-Val-Gln-Met-Pro-Ile-Ser).
- 13 77. The vaccine according to Item 69 wherein the antigenic synthetic peptide
- is 78(Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Gln).
- 78. The vaccine according to Item 69 wherein the antigenic synthetic peptide
- is 87(Gln-Val-Leu-Ser-Thr-thr-Ala-Lys-Glu-Phe).
- 17 79. The vaccine according to claim 69 wherein the antigenic synthetic peptite
- 18 is 126(Ala-Gly-Thr-Ala-pro-Thr-Ala-Gly-Asn-Tyr).

•	80. The vaccine according to Item 69 wherein the antigenic synthetic peptide
1	is 133(Gly-Asn-Tyr-Ser-Gly-Val-Val-Ser-Leu-Val).
2	
_	81. The vaccine according to Item 67 wherein the antigenic synthetic peptide
3	is selected from the group consisting essentially of 3(Lys-Ana-Ile-Thr-Val-Thr-
4	•
5	Ala-Ser-Val), 11(Val-Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
6	22(Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
7	22(Gly-Asn-Ala-Leu-170 Bet 190
8	
9	Glu-Ser-Tyr-Arg-Val),
10	32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe),
11	38(Lys-Thr-Phe-Glu-Ser-Tyr-Arg-Val),
12	66(Pro-Gln-Leu-Thr-Asp-Val-Leu-Asn-Ser),
13	93(Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala),
14	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr),
15	127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
16	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-
17	Tyr-Ser), and mixtures thereof.
	82. The vaccine according to Item 81 wherein the antigenic synthetic peptide
18	or an Ana He-Thr-Val-Thr-Ala-Ser-Val).
19	83. The vaccine according to Item 81 wherein the antigenic synthetic peptide
20	is 11(Val-Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp).
21	is 11(Val-Asp-Pro-Val-16 1 applied 1) is 11(Val-Asp-Pro-Val-16 1 applied 2). The vaccine according to Item 81 wherein the antigenic synthetic peptide
22	
23	is 22(Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val). 85. The vaccine according to Item 81 wherein the antigenic synthetic peptide
24	85. The vaccine according to Item of whateau are The Phe-Glu-Ser-Tyr-Arg-
25	is 32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-Thr-Phe-Glu-Ser-Tyr-Arg-
26	Val).

	-64-
1	86. The vaccine according to Item 81 wherein the antigenic synthetic peptide
2	Sar Pro-Ala-Ser-Lys-Thr-Phe).
3	87 The vaccine according to Item 81 wherein the antigenic synthetic peptide
4	The Clu-Ser-Tvr-Arg-Val).
5	is 38(Lys-Thr-Phe-Gla-Sea 1). 88. The vaccine according to Item 81 wherein the antigenic synthetic peptide
6	Cla Law Thr-Asn-Val-Leu-Asn-Ser).
7	is 66(Pro-Gin-Leu-Fin 715) 89. The vaccine according to Item 81 wherein the antigenic synthetic peptide
8	Glu-Phe-Glu-Ala-Ala-Ala).
9	is 93(Ala-Lys-Glu-ric Gla-ric 93). The vaccine according to Item 81 wherein the antigenic synthetic peptide
10	Ala-Gly-Thr-Ala-Pro-Thr).
11	91. The vaccine according to Item 82 wherein the antigenic synthetic peptide
12	is 127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser).
13	
14	92. The vaccine according to Item 82 wherein the antigenic synthetic peptide
15	Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-1 yr-Ser).
16	is 124(Lys-1nt-Ala-Cs) 124 93. The vaccine according to Item 67 wherein the antigenic synthetic peptide 93. As The Val-Thr-Val-Thr-
17	93. The vaccine according to an issue selected from the group consisting essentially of 3(Lys-Asn-Ile-Thr-Val-Thr-
18 ·	Ala-Ser-Bal-Asp-Pro),
19	8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gin-Ala-Asp),
20	11(Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gin-Ala-Asp),
21	20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
22	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
23	124(Lys-Thr-Ala-Gly-Ala-Gly-Asn-Tyr-Ser), and mixtures thereof. 126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and mixtures thereof.
24	126(Ala-Gly-1nr-Ala-110 110) 126(Ala-Gly-1nr-Ala-110 110) 94. The vaccine according to Item 93 wherein the antigenic synthetic peptide
25	is 3(Lys-Asn-Ile-Thr-Val-Thr-Ala-Ser-Bal-Asp-Pro).
26	95. The vaccine according to Item 93 wherein the antigenic synthetic peptide
27	is 8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-Ile-Asp-LeuLeu-Gln-Ala-Asp).

1	96. The vaccine according to Item 93 wherein the antigenic synthetic peptide
2	ic 11 (Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gin-ala-Asp).
3	97. The vaccine according to Item 93 wherein the antigenic synthetic peptide
4	ic 20(Ala-Asp-Gly-Asp-Ala-Leu-Pro-Ser-Ala-Val).
5	98. The vaccine according to Item 93 wherein the antigenic synthetic peptide
6	in 1240 vs-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser).
7	99. The vaccine according to Item 93 wherein the antigenic synthetic peptitie
8	in 126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser).
9	100. The method of Item 54, wherein the viral infection is caused by a virus
10	selected from the group consisting essentially of hepatitis A, hepatitis B,
11	henatitis C. Varicella-Zoster virus, Epstein-Barr virus, Rotaviruses, polio
12	virus human immunodeficiency virus (HIV), herpes simplex virus type 1,
13	human retroviruses, herpes simplex virus type 2, Ebola virus, cytomegalo
14	viruses. Herres Simplex viruses, Human cytomegalovirus, Varicella-Zostei
15	Virus, Enstein-Barr Virus, Poxvirus, Influenza viruses, Parainfluenza viruses,
16	Respiratory Syncytial virus, Rhinoviruses, Coronaviruses, Adenoviruses,
17	Measles virus, Mumps virus, Robella Virus, Human Parvoviruses,
18	A theories Rabies virus, Enteroviruses, reoviruses, Viruses Causing
19	costmenteritis Henatitis Viruses, Filoviruses, Arenaaviruses, Papulomaviruses,
20	Polyomaviruses, Human Immunodeficiency viruses, Human Retroviruses, and
	Speciform Encephalopathies.
21	101. The method in accordance with Item 49 comprising administering orang
22	to said mammal an immunogenic amount of a pharmaceutical composition
23	consisting essentially of an antigen in the amount of .1 to 1%.
24	A rescine for the immunization of a mammal against infection by
25	the respice organisms consisting essentially of an antigen in the amount of the
26	to 1% encapsulated within a biodegradable-biocompatible polymeric poly(DL-
27	W 170 Cilcapsonana

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	the relumns is end-capped or a blend of
1	lactide-co-glycolide) matrix wherein the polymer is end-capped or a blend of
2	uncapped and end-capped polymers.
3	103. The vaccine according to Item 100 wherein the polymer is a blend of
4	end-capped and uncapped polymers.
5	104. The vaccine according to Item 103 wherein the relative ratio between the
6	to side and alreadide component is within the range of 90/10 to 40/00.
7	The vaccine according to Item 103 wherein the relative ratio between the
8	amount of lactide and glycolide component is within the range of 48/52 to
9	52/48. 106. The vaccine according to Item 102 wherein the antigen is a bacteria or
10	
11	derivatives thereof. 107. The vaccine according to Item 103 wherein the antigen is a virus or
12	107. The vaccine according to Item 105 mass
13	derivatives thereof.
14	108. The vaccine according to Item 103 wherein the antigens is a parasite or
15	derivative thereof.
16	109. The vaccine according to Item 103 wherein the antigen is a fungus or
17	derivative thereof.
18	110. The vaccine according to Item 106 wherein the bacteria is selected from
19	secretaring essentially of Salmonella typhi, Shigella Soulier, Shigella
20	Shipella dysenteriae, Shigella boydii, Eschena coll, Viono chorest
21	B. 2. Group E. Group G, Group I, Group I, Listeria, Erystetoutical
22	Aerobic pathogenic Actinomycetales, Enterobacteriaceae,
	Vibrio aeromonas, Plesiomonas, Helicobacter, W. succinogenes, Acmess
23	bacter spp., Foavobacterium, Pseudomonas, Legionella, Brucella,
24	Haemophilus, Bordetalla, Mycoplasmas, Gardnerella, Streptobacillus,
25	Haemophilus, Boruelana, Marie
26	Spirillum, Calymmatobacterium, Cioasses, and Cocci, Leptospira, Anaerobic Gram-negative Bacteria including bacilli and Cocci,
27	Leptospira, Anaerobic Gram-negative pactoria

	staphylococcus, clostridium, Enteroccus, Streptoccus, Aerococcus,
1	Planococcus, Stomatococcus, Micrococcus, Lactoccus, Germella, Pediococcus,
2	Leuconostoc, Bacillus, Neisseria, Branhamella, Coryne bacterium,
3	campylobacter, Arcanobacterium haemolyticum, Rhodococcus spp.
4	
5	Rhodococcus, Group A-4.
6	111. The vaccine of Item 107 wherein the virus is selected from the group
7	consisting essentially of hepatitis A, hepatitis B, hepatitis C, Varicella-Zoster
8	virus, Epstein-Barr virus, Rotaviruses, polio virus, human immunodeficiency
•	herpes simplex virus type 1, human retroviruses, herpes simplex
9	virus (HV), herpes Virus (HV), herpes Simplex viruses, Human virus type 2, Ebola virus, cytomegalo viruses, Herpes Simplex viruses, Human
10	cytomegalovirus, Varicella-Zoster Virus, Epstein-Barr Virus, Poxvirus,
11	cytomegalovirus, Varicena 220000 Pespiratory Syncytial virus,
12	Influenza viruses, Parainfluenza viruses, Respiratory Syncytial virus,
13	Rhinoviruses, Coronaviruses, Adenoviruses, Measles virus, Mumps virus,
14	Daballa Virus Human Parvoviruses, Arboviruses, Rabies virus, Enteroviruses,
15	recognizates. Viruses Causing gastroenteritis Hepatitis Viruses, Filoviruses,
16	Arenaaviruses, Papillomaviruses, Polyomaviruses, Human Immunodericiency
17	viruses. Human Retroviruses, and Spongiform Encephalopathies.
	112. An immunostimulating composition comprising encapsulating-
18	microspheres, which may contain a pharmaceutically-acceptable adjuvant,
19	microspheres, which may contain a phase between 1 nanogram (ng) to 10
20	wherein said microspheres having a diameter between 1 nanogram (ng) to 10
21	microns (um) are comprised of (a) a biodegradable-biocompatible poly (DL-
22	lactide-co-glycolide) as the bulk matrix, wherein the copolymer (lactide to
23	glycolide L/G) ratio for uncapped and end-capped polymer is 0/100 to 1/99
24	de la compunaçõe de comprising a bacteria, virus, fungus,
25	parasite, or derivative thereof, that serves to elicit the production of antibodies
26	in animal subjects.

113. An immunostimulating composition according to Item 112 wherein the amount of said immunogenic substance is within the range of 0.1 to 1.5% 1 2 based on the volume of said bulk matrix. 114. An immunostimulating composition according to Item 10 wherein the 3 immunogenic substance comprises Colony Factor Antigen (CFA/II), hepatitis B surface antigen (HBsAg), a mixture thereof physiologically similar antigen. 5 6 115. An immunostimulating composition according to Item 113 wherein the relative ratio between the lactide and glycolide component is within the range 7 8 of 48/52 to 52/48. 116. An immunostimulating composition according to Item 113 wherein the 9 size of more than 50% of said microspheres is between 5 to 10 um in diameter 10 11 by volume. 117. An immunostimulating composition according to Item 113 wherein the 12 immunogenic substance is the synthetic peptide representing the peptide 13 fragment beginning with the amino acid residue 63 through 78 of Pilus Protein 14 CS3, said residue having the amino acid sequence, 63(Ser-Lys-Asn-Gly-Thr-15 16 Val-Thr-Try-Ala-His-Glu-Thr-Asn-Asn-Ser-Ala). 118. A vaccine comprising an immunostimulating composition of Item 113 17 18 and a sterile, pharmaceutically-acceptable carrier therefor. 119. A vaccine comprising an immunostimulating composition of Item 118 19 wherein said immunogenic substance is Colony Factor Antigen (CFA/II). 20 120. A vaccine comprising an immunostimulating composition of Item 119 21 wherein said immunogenic substance is hepatitis B surface antigen (HBsAg). 22 121. A method for the vaccination against bacterial infection comprising 23 administering to a human, an antibactericidally effective amount of a 24 25 composition of Item 118. 122. A method according to Item 121 wherein the bacterial infection is caused 26 27

	typhi, Shigella Sonnei, Shigella Flexneri, Shigella dysenteriae, Shigella boyun,
1	Escheria coli, Vibrio cholera, Group D-2, Group E, Group G, Group I, Group
2	Escheria coli, Vibrio cholera, Citono De la Aerobic pathogenic
3	1, Listeria, Erysipelothrix, Mycobacterium, Aerobic pathogenic
4	Actinomycetales, Enterobacteriacese, Vibrio, aeromonas, Plesiomonas,
5	Helicobacter, W. succinogenes, Acineto bacter spp., Foavobacterium,
6	Legionella, Brucella, Haemophilus, Bordezula, Mysser
7	constancially Spirilly, Calymmatopacterium, Stores
	Posselia Leptospira, Anaerobic Gram-negative Datastra
8	Anaerobic gram-Positive Nonsporeior mine Parameter
9	etaphylococcus, clostridium, Enteroccus, Sucotovers,
10	Aerococcus, Planococcus, Stomatococcus, Micrococcus, Lactoccus, Germella,
11	Aerococcus, Planococcus, Prantococcus, Prant
12	Pediococcus, Leuconosioc, Dacines, Leuconosioc, Leuconosioc, Dacines, Leuconosioc, L
13	
14	Rhodococcus, Group A-4.
15	Rhodococcus, Orosa re- Rhodococcus, Orosa re- 123. A method for the vaccination against viral infection comprising
16	123. A method for the vaccination of administering to a human an antivirally effective amount of a composition of
17	Item 108.
18	Item 108. 124. A diagnostic assay for bacterial infections comprising a composition of
19	Item 7.
20	Item 7. 125. A method of preparing an immunotherapeutic agent against infections
21	caused by a bacteria comprising the steps of (1) immunizing a plasma donor
22	with a vaccine according to Item 52 such that a hyperimmune globulin is
23	tich contains antibodies directed against the balleting (2)
	signal and (3) purifying the hyperimmune grounds
24	an immunotherapeutic agent against misses
25	the step of immunizing a plasma dono.
26	vaccine according to Item 126 such that hyperimmune globulin is produced
27	vaccine according to them.

1	127. An immunotherapy method comprising the step of administering to a
2	subject an immunostimulatory amount of hyperimmune globulin prepared
3	according to Item 125.
4	128. An immunotherapy method comprising the step of administering to a
5	subject an immunostimulatory amount of hyperimmune globulin prepared
6	according to Item 125.
7	129. A method for the protection against infection of a subject by
8	enteropathogenic organisms or hepatitis B virus comprising administering to
9	said subject an immunogenic amount of an immunostimulating composition of
10	Item 112.
11	130. A method according to Item 127 wherein the immunostimulating
12	composition is administered orally.
13	131. A method according to Item 127 wherein the immunostimulating
14	composition is administered parenterally.
15	132. A method according to Item 127 wherein the immunostimulating
16	composition is administered in four separate doses on day 0, day 7, day 14,
17	and day 28.
18	133. A method according to Item 114 wherein the immunogenic substance is
19	the synthetic peptide representing the peptide fragment beginning with the
20	amino acid residue 63 through 78 of Pilus Protein CS3 said residue having the
21	amino acid sequence 63(Ser-Lys-Asn-Gly-Thr-Val-Thr-Try-ala-His-Glu-thr-an-
22	Asn-Ser-Ala).
23	134. A method for the protection against or therapeutic treatment of bacterial
24	infection in the soft tissue or bone of a mammal comprising administering
	locally to said mammal a bactericidally-effective amount of a composition of
25	Item 2, wherein the active material is an antibiotic which is controlled release
26	within a period of about 1 to 100 days
27	Midmi of free and a second a second and a second a second and a second

	135. The method according to Item 134 wherein the biodegradable poly(DL-
1	135. The method according to remain a lactide-co-glycolide) is a blend of uncapped and end-capped forms having a lactide-co-glycolide) is a blend of uncapped and end-capped forms having a
2	lactide-co-glycolide) is a blend of uncapped and glycolide component within
3	lactide-co-glycolide) is a blend of each of lactide and glycolide component within relative ratio between the amount of lactide and glycolide component within
4	100
5	the range of 100/0 to 1799. 136. A method according to Item 135 wherein the bacterial infection is (1) a
6	subcutaneous infection secondary to contaminated abdominal surgery, (2) an
7	infection surrounding prosthetic devices and vascular grafts, (3) ocular
8	infection surrounding products of infections, (5) orthopedic infections, including infections, (4) topical skin infections, (5) orthopedic infections, including
9	osteomyelitis, and (6) oral infections.
10	osteomyelitis, and (6) or a large of the oral infections are 137. The method according to Item 136 wherein the oral infections are
11	· 1-pel dicesse.
12	pericoronitis or periodontal discussions. 138. The method according to Item 135 wherein the administration is effected
13	prior to infection.
14	prior to infection. 139. The method according to Item 135 wherein the administration is effected
15	subsequent to infection.
16	subsequent to infection. 140. The method according to Item 135 wherein said animal is a human. 141. The method according to Item 135 wherein said animal is a nonhuman.
17	141. The method according to Item 135 whose sometising applying to the soft
18	141. The method accordance with Item 135 comprising applying to the soft 142. The method in accordance with Item 135 comprising applying to the soft
19	142. The method in accordance tissue or bone tissue of said animal a bactericidally-effective amount of a tissue or bone tissue of said animal a bactericidally-effective amount of a
20	pharmaceutical composition consisting essentially of an antibiotic in the ant,
21	pharmaceutical compositions of a beta-lactam, aminoglycolide, selected from the group consisting of a beta-lactam, aminoglycolide,
22	selected from the group consistency selected from the group consistency polymyxin-b, Amphotericin B, Aztreonam, cephalosporins, chloramphenical, polymyxin-b, Amphotericin B, Aztreonam, cephalosporins, chloramphenical,
23	polymyxin-b, Amphotester-by fusidans, lincosamides, macrolides, methronidazole, nitro-furation, fusidans, lincosamides, macrolides, methronidazole, nitro-furation,
24	fusidans, lincosamides, macronees, fusidans, lincosamides,
25	eicoplanin, imidazoles, and differences
26	damedable poly(DL-izcude-or gr)
27	wherein the amount of the lactide and glycolide (20)

rix which is present in the amount of from 40 to 95 percent, resulting in
s the enid antihiotic OVET a
controlled release of a bacteriacidal amount of the said antibiotic over a
and of from 1 to 100 days.
2. The method of Item 142 wherein the polymeric matrix consists
wherein the relative ratio
tween the amount of lactide and glycolide (L/G) component is within the
5 49/52 to 52/48.
weeked of Item 142 wherein the bacterial infection is caused by a
the second registant bacteria selected from the group consisting essentially
of Enterobacteriaceae; Klebsiella sp.; Bacteroides sp. Enterococci; Proteus sp.;
Streptococcus sp.; Staphylococcus sp.; Pseudomonas sp.; Neisseria sp.;
Pedptostreptococcus sp.; Fusobacterium sp.; Actinomyces sp.; Mycobacterium
Sp.; Listeria sp.; Corynebacterium sp.; Proprionibacterium sp.; Actinobacillus
sp.; Listeria sp.; Corynebacter sp.; Campylobacter sp.; cytophaga sp.; sp.; Aerobacter sp.; Borrelia sp.; Campylobacter sp.; cytophaga sp.;
Sp.; Aerobacter sp.; Borrella sp., Santonacter aerogenes, Peptococcus sp., Pasteurella sp.; Clostridium sp., Enterobacter aerogenes, Peptococcus
Proteus vulgaris, Proteus morganii, Staphylococcus aureus, Smeptococcus
Proteus vulgaris, Proteus morganii, Staphyrocept and Legionella
pyogenes, Actinomyces sp., Campylobacter fetus, and Legionella
progenes, Actinomyses and progenes, Actinomyses and methicillin-resistant pneumophila, ampillin-resistant strain of S. aureus, and methicillin-resistant
strain of S. aureus.
strain of S. aureus. 145. The method of Item 142 wherein the antibiotic is selected from the group
consisting essentially of a beta-lactam, aminoglycolide, polymyxin-B,
amphotericin B, aztreonam, cephalosporins, chloramphenicol, fusidans,
amphotericin B, azziconam, or lincosamides, macrolides, methronidazole, nitro-furantoin, Imipenem/cilastin,
quinolones, rifampin, polyenes, tetracycline, sulfonamides, unicaroptan,
in anicoplania imidazoles, and erythromycin.
The method of Item 145 wherein the beta-lactam is cepinalesperation
146. The method of Item 145 wherein the beta-lactam is penicillin.

	· · · · · · · · · · · · · · · · · · ·
1	149. The method of Item 145 wherein the aminoglycolide is amikacin.
2	as a method of Item 145 wherein the aminoglycolide is toolainyem.
3	The method of Item 145 wherein the aminoglycolide is animyoun
	152. The method of Item 145 wherein the beta-lactam is an amplement
5	153. The method of Item 152 wherein the polymeric matrix consists
	wherein the relative ratio
6 7	between the amount of lactide and glycolide (L/G) component is within the
	C 48/52 to 58/42.
8	The method of Item 152 wherein the ampicillin is present in an amount
9	of from 5 to 60 percent and the amount of polymeric matrix is from 40 to 95
10	3-3
11	percent. 155. The process of using the composition of Item 1 to treat humans in need,
12	thereof, suffering from diseases and/or ailments from the group consisting of:
13	viral infections; bacterial infections; fungal infections; parastic infections and
14	more specific diseases and/or ailments; such as as, aids; alzheimer's dementia;
15	angiogenesis diseases; aphthour ulcers in AIDS patients; asthma; atopic
16	angiogenesis diseases; aphiliotic erectiona; benign prostatic hypertrophy; blood dermatitis; psoriasis; basal cell carcinoma; benign prostatic hypertrophy; blood
17	dermatitis; psoriasis; basar cen earestenay substitute; blood substitute in surgery patients; blood substitute in trauma
18	substitute; blood substitute in surgery parameter, substitute; blood substitute in surgery parameter, metastatic; cachexia in patients; breast cancer; breast cancer; cutaneous & metastatic; cachexia in
19	patients; breast cancer; breast cancer; premonia; sexually transmitted
20	AIDS; campylobacter infection; cancer; pnemonia; sexually transmitted diseases (STDs); cancer; viral dieases; candida albicians in AIDS and cancer;
21	diseases (STDs); cancer; viral dieases, canalism and cancer; parkinson's
22	candidiasis in HIV infection; pain in cancer; pancreatic cancer; parkinson's
23	disease; peritumoral brain edema; postoperative adhesions (prevent);
24	proliferative diseases; prostate cancer; ragweed allergy; renal disease;
25	restenosis; rheumatoid arthritis; rheumatoid arthritis; allergies; rotavirus
26	infection; scalp psoriasis; septic shock; small-cell lung cancer; solid tumors;
27	stroke; thrombosis; type I diabetes; type I diabetes w/kidney transplants: type

1	rthythm disorders; central nervous system diseases; central nervous system
	ticardess: cervical dystonia (spasmodic torticollis); choridal neovascularization,
2	chronic hepatitis c, b and a; colitis associated with antibiotics; colorectal
3	corporary artery thrombosis; cryptosporidiosis in AIDS;
4	cryptosporidium parvum diarrhea in AIDS; cystic fibrosis; cytomegalovirus
5	disease; depression; social phobias; panic disorder; diabetic complications;
6	disease; depression, social process, in disabetic eye disease; diarrhea associated with antibiotics; erectile dysfunction;
7	disabetic eye disease; diarrilea association disease in transplant patients; growth hormone genital herpes; graft-vs host disease in transplant patients; growth hormone
8	genital herpes; graft-vs host disease in transport p
9	deficiency; head and neck cancer; head trauma; stroke; heparin neutralization
10	after cardiac bypass; hepatocellular carcinoma; HIV; HIV infection;
11	huntington's disease; CNS diseases; hypercholesterolemia; hypertension;
12	inflammation; inflammation and angiogensis; inflammation in cardiopulmonary
13	bypass; influenza; migrain head ache; interstitial cystitis; kaposi's sarcoma;
14	in AIDS; lung cancer; melanoma; molluscum contagiosum ai
	a visible sclerosis; neoplastic meningitis from solid rumors, non-small
15	transplant rejection; osteoarthritis; meumatoid addition,
16	osteoporosis; drug addiction; shock; ovarian cancer; Amebiasis; Babesiasis;
17	Cryptosporidiosis; Cysticercosis,
18	Chagas' disease (Typanococcosis; Giardiasis; Leishmaniasis; Malaria; Fascioliasis; Filariasis; Echinococcosis; Giardiasis; Leishmaniasis; Malaria;
19	Fascioliasis; Filariasis; Etimiocescopi, Paragonimiasis; Pneumocystosis; Schistosomiasis; Strongylodiasis;
20	Paragonimiasis; Pneumocysiosis, Schabbook Toxocariasis; Toxoplasmosis; Trichinellosis; Trichomoniasis; yeast infection;
21	Toxocariasis; Toxoplasmosis; Inclinicatosas,
22	and pain.
23	and pain. 156. A vaccine for prepared from the composition of Item 1 to prevent the
24	occurence in humans of diseases and/or ailments comprising viral infections;
25	infections; parastic infections and more specific
26	liances and/or ailments: such as as, aids; alzheimer's dementa, angrogeniere
27	diseases; aphthour ulcers in AIDS patients; asthma; atopic dermatitis;
21	· · · · · · · · · · · · · · · · · · ·

	-/J-
1	blood substitute in surgery patients; blood substitute in trauma patients; breast
	hand cancer: cutaneous & metastatic; cachexia in Aids,
2	remove the content infection; cancer; pnemonia; sexually transmitted discard
3	simil dieses; candida albicians in AIDS and Caned,
4	(STDs); cancer; vital detection; pain in cancer; pancreatic cancer; parkinson's candidiasis in HIV infection; pain in cancer; pancreatic cancer; parkinson's
5	disease; peritumoral brain edema; postoperative adhesions (prevent);
6	proliferative diseases; prostate cancer; ragweed allergy; renal disease;
7	restenosis; rheumatoid arthritis; rheumatoid arthritis; allergies; rotavirus
8	infection; scalp psoriasis; septic shock; small-cell lung cancer; solid tumors;
9	infection; scalp psoriasis; septic sheet, stroke; thrombosis; type I diabetes; type I diabetes w/kidney transplants; type stroke; thrombosis; type I diabetes; type I diabetes w/kidney transplants; type
10	stroke; thrombosis; type I diabetes, type I diabetes, type I diabetes; typ
11	II diabetes; viseral leishmaniasis, mania, property in diabetes; viseral leishmania, mania, property in diabetes; viseral
12	rthythm disorders; central nervous system electrical recovery disorders; cervical dystonia (spasmodic torticollis); choridal neovascularization;
13	disorders; cervical dystonia (spasmoule with antibiotics; colorectal
14	chronic hepatitis c, b and a; colitis associated with antibiotics; colorectal
15	cancer; coronary artery thrombosis; cryptosporidiosis in AIDS;
16	cancer; coronaly altery diagrams in AIDS; cystic fibrosis; cytomegalovirus cryptosporidium parvum diarrhea in AIDS; cystic fibrosis; cytomegalovirus
17	disease; depression; social phobias; panic disorder; diabetic complications;
18	disease; depression, seems procession, disease; depression, seems process; disease; disease; diarrhea associated with antibiotics; erectile dysfunction;
19	genital herpes; graft-vs host disease in transplant patients; growth hormone
20	deficiency; head and neck cancer; head trauma; stroke; heparin neutralization
21	lice hypass: henatocellular carcinoma; HIV; HIV infection,
22	CNS diseases; hypercholesterolemia; nyperchasson,
23	inflammation and angiogensis; inflammation in cardiopersis
24	migrain head ache; interstitial cystins; kaposi s successions
25	in ATDS: lung cancer; melanoma; monuscum contragram
26	and a colomois: neoplastic meningitis from solid tunions, non-
	AIDS; multiple scierosis, inceptual AIDS; multiple scierosis, inceptual arthritis; cell lung cancer; organ transplant rejection; osteoarthritis; rheumatoid arthritis;
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	Chagas' disease (Trypanosoma cruzi); Cryptosporidiosis; Cysticercosis;
1	Chagas' disease (Trypanosonia or Chagas') disease (Trypanosonia or Chagas' disease (Trypanosonia or Chagas' disease (Trypanosonia or Chagas' disease (Trypanosonia or Chagas') disease (Trypanosonia or Chagas' disease (Trypanosonia or Chagas' disease (Trypanosonia or Chagas' disease (Trypanosonia o
2	Fascioliasis; Filariasis; Echinococcas, Fascioliasis; Strongylodiasis;
3	Paragonimiasis; Pneumocystosis; Schistosomiasis; Strongylodiasis; Paragonimiasis; Pneumocystosis; Schistosomiasis; Strongylodiasis;
4	Paragonimiasis; Phetimocysosas, or Trichinellosis; Trichomoniasis; yeast infection; Toxocariasis; Toxoplasmosis; Trichinellosis; Trichomoniasis; yeast infection;
5	and pain.
6	and pain. As noted, in the Summary of the Invention section herein, a discussion
	of this invention will be presented as Phases I, II and III.
7	PHASE I
8	This illustrative phase of the inventionpresents the novel use of a
9	This illustrative phase of the
10	pharmaceutical composition, a micro of master a biodegradable polymeric
11	which comprises an antibiotic encapsulated within a biodegradable polymeric
12	DI lactide-co-glycolide) (DL-PLG) in all of the
	and mammals to prevent bacterial infections and the
13	disting humans and non-human mammals) with
14	of mammals (including finance) infections. Microcapsules and microspheres are usually powders consisting of
15	of 2 millimeter or less in diameter, usually 500 me
16	or less in diameter. If the particles are less than 1 micron, they are often
17	or less in diameter. If the particles are less the most part, the difference
18	or less in diameter. If the parties or nanospheres. For the most part, the difference referred to as nanocapsules or nanospheres. For the most part, the difference
19	is their size, titel meaning is their size, titel meaning
	Similarly, the difference between microsphiese
20	their size: their internal structure is about the same.
21	A microcapsule (or nanocapsule) has its encapsulated material,
22	A microcapsule (of interest within a unique membrane,
23	herein after referred to as agent, centrally located within a unique membrane,
24	usually a polymeric membrane. This membrane may be termed a
	and is usually a polyment matthm.
25	nemente microcapsules designed for conduction
26	is since selegge their agent at a constant rate (zero-order
27	applications release

Hereinaster, the term microcapsule will include nanocapsules, and particles in general that comprise a central core surrounded by a unique outer membrane. 1 2 A microsphere has its agent dispersed throughout the particle; that is, the internal structure is a matrix of the agent and excipient, usually a 3 polymer excipient. Usually controlled-release microspheres release their agent 4 at a declining rate (first-order). But microspheres can be designed to release 5 agents at a near zero-order rate. Microspheres tend to be more difficult to 6 7 rupture as compared to microcapsules because their internal structure is 8 stronger. Hereinafter, the term microspheres will include nanospheres, 9 microparticles, nanoparticles, microsponges (porous microspheres) and particles in general, with an internal structure comprising a matrix of agent 10 11 and excipient. 12 One can use other terms to describe larger microcapsules or microspheres, that is, particles greater than 500 micrometer to 7 millimeter or 13 14 larger. These terms are macrocapsules, macrospheres, macrobeads and 15 beads. Macrocapsules, macrospheres, macrobeads and beads will be used 16 interchangably herein. 17 More particularly, the applicants have discovered efficacious pharmaceutical compositions wherein the relative amounts of antibiotic to the 18 polymer matrix are within the ranges of 5 to 60 preferred that relative ratio 19 20 between the lactide and glycolide component of the poly(DL-lactide-co-glycolide) is within the range of 40:60 to 100:0, most 21 preferably. Applicants' most preferred composition consists essentially of 30 22 to 40(core loading) and 60 to 70 poly(DL-lactide-co-glycolide) (DL-PLG). 23 However, it is understood that effective core loads for other antibiotics will be 24 influenced by the nature of the drug, the microbialetiology and type of 25 infection being prevented and/or treated. From a biological perspective, the 26 27

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under physiologic conditions to products that are nontoxic and readily metabolized. Similar polymeric compositions which afford in vitro release kinetics, as discussed below for DL-PLG, are considered by applicants to be within the scope of this invention. Applicants have discovered that antibiotic encapsulated microcapsules/spheres or macrocapsules/spheres (beads) having a diameter within the range of about 40 microns to about 7 millimeters to be especially useful in the practice of this invention.

Surprisingly, applicants have discovered an extremely effective method of treating bacterial infections of soft-tissue or (bone osteomyelitis) and preventing these type infections with antibiotics such as beta-lactams, aminoglycosides, polymyxin-B, amphotericin B, aztreonam, cephalosporins, chloramphenicol, fusidans, lincosamides, macrolides, metronidazole, nitro-furantion, Imipenem/cilastin, quinolones, rifampin, polyenes, tetracycline, sulfonamides, trimethoprim, vancomycin, teicoplanin, imidazoles, and erythromycin 1) micro- and macroencapsulated or 2) microand macrospheres formulated within a polymeric matrix such as a poly(DL-lactide-co-glycolide), which has been formulated to release the antibiotic at a controlled, programmed rate over a desirable extended period of time. The microcapsules/spheres have been found to be effective when applied locally, including topically, to open contaminated wounds thereby facilitating the release of the antibiotic from multiple sites within the tissue in a manner which concentrates the antibiotic in the area of need. Similarly, the encapsulated antibiotics of this invention both in the microcapsule/sphere and macrocapsule/sphere (bead) form are effective for the prevention and treatment of orthopedic infections that include osteomyelitis, contaminated open fractures, and exchange revision arthroplasty. The macrocapsules/sphere

addition the option to the surgeon of using the subject invention as a packing material for dead space. The subject invention offers an optimal treatment for orthopaedic infections because release of the antibiotic from the micro- or macrocapsule/sphere is completely controllable over time; antibiotic can be encapsulated into the sphere; the sphere can be made of any size; and unlike the methylmethracrylate beads, the subject invention biodegrades over time to nontoxic products and does not have to be surgically removed from the treated site. Since virtually any antibiotic can be encapsulated into the polymer the instant invention can be used to sustain release all known antibiotics.

Applicants have discovered and/or contemplate that local application of microencapsulated or macroencapsulated antibiotic provides immediate, direct, and sustained dosing which targets the antibiotic to the pre-or post infected soft-tissue or bone site, and minimizes problems inherent in systemic drug administration. It appears to applicants that there is a significant reduction of nonspecific binding of antibiotic to body proteins, while in route to targeted sites when the antibiotic has been encapsulated in accordance with this invention. Additionally, antibiotics with short half-lives can be used more efficiently, undesirable side-effects can be minimized, and multiple dosing can be eliminated. These attributes satisfy a long-felt need to improve the effectiveness and predictability of drug delivery to accomplish the desired clinical result in patients.

The ability to concentrate the antibiotic within the wound site ensures an extended period of direct contact between an effective antibiotic level and the infecting microorganisms. Many drugs have a therapeutic range below which they are ineffective and above which they are toxic. Oscillating drug levels, commonly observed following systemic administration, may cause alternating periods of ineffectiveness and toxicity. A single dose of

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-80desired therapeutic range. Applicants have discovered that microencapsulated or macroencapsulated heavy concentrated doses of antibiotics are effective for 1 the treatment and prevention of infections caused by antibiotic-resistant 2 3 bacteria. 4 Topical application of the antibiotic microcapsule/ sphere formulation to infected wounds allows local application of the antibiotic in a 5 single dose, whereby an initial burst of antibiotic for immediate soft- or 6 hard-tissue perfusion, followed by a prolonged, effective level of antibiotic is 7 achieved in the tissue at the wound site. Applicants contemplate herein 8 antibiotic microcapsules/spheres and macrocapsules/spheres consisting of an 9 antibiotic and DL-PLG and the summarized results of illustrative experiments 10 11 that evaluated the prototype microcapsules in vitro and in vivo. The subject invention is successful in preventing and treating 12 (1) soft-tissue infections, (2) osteomyelitis, and (3) infections surrounding 13 14 internally fixed fractures. These results were confirmed using the 15 microcapsule/sphere form of the encapsulated antibiotics. The microcapsule/sphere and macrocapsule/sphere are also of value in numerous 16 other applications including soft-tissue infections that involve, but are not 17 limited to the prevention and treatment of (1) subcutaneous infections 18 secondary to contaminated abdominal surgery, (2) infections surrounding 19 prosthetic devices and vascular grafts, (3) ocular infections, (4) topical skin 20 infections, and (5) in oral infections such as pericoronitis and periodontal 21 22 disease. The biodegradation rate of the excipient is controllable because 23 it is related to the mole ratio of the constituent monomers, the excipient 24 molecular weight and the surface area of the microcapsules produced. 25

Microcapsules/spheres with diameters of 250 micrometers or less are

	aerosol spray. The macrocapsules/spheres are manually placed in the tissue
1	aerosol spray. The macrocapsules spray
2	on bone by the surgeon at the time of surgical debridement. Due to the
	in a sharmacokinetic advantages realized with the continuous delivery of
3	antibiotic into tissue from a controlled-release vehicle, applicants have found
4	antibiotic into tissue from a controlled the antibiotic into tissue from the antibiotic int
5	that a small total dose is required to obtain an optimal therapeutic effect.
	VII. EXAMPLES
6 .	The herein offered examples provide methods for illustrating,
7	The never of this invention in the treatment
8	without any implied limitation, the practice of this invention in the treatment
9	of bacterial wound infections.
	The profile of the representative experiments have been chosen
10	to illustrate the antibacterial activity of antibiotic-polymeric matrix
11	to illustrate the antibacterial activity of
12	composites.
13	All temperatures not otherwise indicated are in degrees Celcius
13	(°C) and parts or percentages are given by weight.
14	(°C) and parts of percentage
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MATERIALS AND METHODS

A. Microcapsules/spheres. The ampicillin anhydrate microspheres used in these studies (Composite Batch D 856-038-1) consisted of 30.7 wt in a copolymer of 52:48 poly (DL-lactideco-glycolide). The size of the microspheres ranged from 45 to 150 microns and they were sterilized with 2.0 Mrad of gamma irradiation.

Animals. New Zealand white rabbits (Dutchland Laboratories, Denver, Pa.), weighing 2.0 to 2.5 kg each, were used. The animals were housed in individual cages and were fed a standard laboratory diet. The experiments described herein were conducted in accordance with the principles set forth in the Guide for the Care and Use of Laboratory Animals.

EXAMPLE 1

Osteomyelitis Model. The technique used to produce osteomyelitis was a modification of the procedure described previously by Norden. Briefly, New Zealand white rabbits (2.0 - 2.5 kg, each) were anesthetized with ketamine hydrochloride and xylazine and access to the medullary canal was gained by inserting an 18-guage Osgood needle (Betton Dickinson Corp., Rutherford, NI) into the right proximal tibial metaphysis. Through this needle was injected 0.1 ml of 5 Pharmaceuticals, Tenafly, M) followed by injection of approximately 5 x 10° CFU of S. aureus ATCC 6538P.

- The hole in the bone was sealed with bone wax and each animal
- received a single subcutaneous injection of 3-ml TORBUTROLTM (A. ı
- Hunt Valley, MD) for postoperative pain control. 2
- Antibiotic therapy was then initiated either immediately or was J. Buck, 3
- delayed for 7-days as described in detail below. 5

EXAMPLE 2

- The initial experiment Immediate Antibiotic Therapy. 6
- was designed to evaluate the efficacy of local therapy with 7
- microencapsulated ampicillin for the prevention of experimental 8 9
- osteomyelitis. A total of 31 rabbits were infected in the right
- proximal tibia with sodium morrhuate and \underline{S} . aureus and treatment 10 11
- was initiated immediately as follows: 12
- Group A (n = 6) received three daily subcutaneous
- ampicillin sodium 13 aqueous mg/kg/day) of
- (Polycillin- N^{TM} , Bristol Laboratories, Syracuse, NY) at 8-hour 14 15
- intervals for 14 consecutive days;} ló
- Group B (n = 7) received a single intramedullary
- injection of 100 mg of microencapsulated ampicillin containing an 17 18
- equivalent of 30.7 mg of ampicillin anhydrate. The microcapsules/
- spheres were suspended in 0.2-ml of 2injection vehicle) and were 19
- then injected into the medullary canal through the same needle 20
- that was used to inject the sclerosing agent and bacteria; 21 22

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<u>Group C</u> (n = 4) received a single intramedullary injection of 0.12 ml (30.7 mg) of aqueous sodium ampicillin 1 (representing the unencapsulated free drug); and 2 3

Groups D, E, and F (n = 14) served as controls and intramedullary injection of 4 microcapsules (100 mg) without antibiotic; injection vehicle (0.2 either an 5 6 ml) without antibiotic; or no treatment. 7

The animals were observed for a total of 8-weeks during time roentgenograms were obtained to evaluate 8 progression of the disease. All surviving animals were euthanized 9 intraveneously at two months postinfection with T-61 euthanasia 10 solution (1 mg/kg/iv) and the tibiae were harvested for 11 bacteriological analysis as described below. 12 13

EXAMPLE 3

Delayed Antibiotic Therapy Without Debridement. In the 14 second experiment, a total of 30 rabbits were injected in the 15 right proximal tibia with sodium morrhuate and \underline{S} . aureus and the 16 infection was allowed to become established for 7-days. Om Day 7, 17 the animals were reanesthetized and an incision was made over the 18 patellar tendon to expose the tibial tuberosity. A 5-mm drill 19 hole was made in the tibial tuberosity and a trocar, measuring 20 approximately 15 centimeters in length, was inserted into the 21 medullary canal to obtain a marrow specimen for culture. 22 animals were then randomly assigned to the following treatment 23 24

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- groups: 1
- Group A (n = 8) received three daily subcutaneous
- injections of aqueous sodium ampicillin (75mg/kg/day) at 8-hour 2 3
- intervals for 14-days;
- Group B (n = 8) received an intramedullary application 5
- of 150 mg of microencapsulated ampicillin containing an equivalent
- 6 of 45 mg of ampicillin anhydrate. 7
- initially suspended in 0.2 ml of the injection vehicle and then
- aspirated into a sterile trocar. The trocar was then inserted 8
- into the medullary canal through the drill hole in the tibial 10
- tuberosity; 11
- <u>Group C</u> (n = 8) received an intramedullary application
- of 0.18 ml (45 mg) of aqueous sodium ampicillin which was also 12 13
- delivered into the canal with a trocar; and 14
- Group D (n = 6) served as controls and received no 15
- treatment. 16
- Following the implantation of the antibiotics into the 17
- medullary canal, the hole in the tibial tuberosity was sealed with
- bone wax and the incision site was closed with 3-0 Dexon sutures. 19
- All of the surviving animals were euthanized 8 weeks following the 20
- initiation of treatment and the tibiae were harvested for 21

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bacteriological analysis.

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EXAMPLE 4

Delayed Antibiotic Therapy With Debridement. standard treatment of chronic osteomyelitis requires the surgical removal of devitalized osseous tissue, the objective of this experiment was to evaluate the efficacy of local antibiotic 5 therapy with microencapsulated ampicillin anhydrate when used in 6 conjunction with debridement. A total of 30 rabbits were injected 7 in the right proximal tibia with sodium morrhuate and \underline{S} . aureus 8 and the infection was allowed to establish for 7 days. On Day 7 9 each animal underwent a standardized surgical debridement of the 10 infected tibia. The animals were anesthetized and an incision was 11 made to expose the medial aspect of the tibia. A Hall drill was 12 used to decorticate approximately one-third of the bone thereby 13 creating a channel that extended the length of the bone. 14 canal was thoroughly debrided with a curette and then irrigated 15 with 20 ml of sterile saline. Cultures of the marrow were 16 obtained at this time for bacteriological analysis. 17 following completion of the debridement procedure, the animals 18 were randomly assigned to the following treatment groups: 19 20

Group A (n = 10) received 150 mg of microencapsulated ampicillin containing an equivalent of 45 mg of 21 The microcapsules were suspended in 0.2-ml 22 injection vehicle and were then implanted into the debrided canal anhydrate. 23 24

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with a sterile trocar; 1

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Group B (n = 10) received 45 mg of unencapsulated sodium ampicillin in powder form which was applied uniformly into the 2 3 Group C (n = 5) and Group D (n = 5) served as controls debrided canal; and 4 and received either an intramedullary application of placebo 5 microcapsules (150 mg) without antibiotic or (2) an injection 6 vehicle (0.2 ml) without antibiotic, respectively. 7

Immediately following the implantation of the materials into the medullary canal, the incision site was closed with 3-0 9 Dexon sutures and each animal received 3-ml of TorbutrolTM for 310 The animals were consecutive days for postoperative pain. 11 euthanized at 8 weeks following the initiation of treatment and 12 the tibiae were harvested for bacteriological evaluation. 13 14

EXAMPLE 5

Radiographs of the 15 Roentgenographic Evaluation. infected tibiae were obtained at various time intervals and were 16 evaluated by a board certified skeletal radiologist (LMM) using a 17 grading system that was originally developed by Norden et al. 18 Four radiographic parameters (sequestrum formation, periosteal 19 reaction, bone destruction, and extent of disease) were evaluated 20 for each animal and a numerical value was assigned for each 21 variable. The scores were then totaled to arrive at an overall 22 23

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The highest total score possible radiographic severity score. with this grading scheme was +7 and reflected the maximum degree l of radiographic severity. 3

EXAMPLE 6

For bacteriological evaluation, the 4 Cultures of Bone. tibiae were dissected free of adherent soft-tissue and the surface 5 of the bone was cleaned with alcohol pads. The bone was then 6 weighed and crushed to small pieces with a sterile mortar and pestle. The crushed bone was suspended in 5 ml of sterile saline 8 and serial 10-fold dilutions were prepared in 0.1 Each dilution 9 (0.1 ml) was then streaked onto both sheep blood agar and mannitol 10 salt agar plates which were incubated aerobically at 37℃ for 24 11 hours. The recovery of any \underline{s} . aureus colonies from the bones was 12 evidence of a persistent osseous infection and was considered as 13 14 a treatment failure. 15

EXAMPLE 7

Measurement of Serum Ampicillin Levels. 16 experiment where local antibiotic therapy was used in conjunction 17 . with debridement, serum levels of ampicillin were measured for all 18 of the animals treated with either an intramedullary application 19 microencapsulated ampicillin anhydrate 20 unencapsulated free drug (Group B). Serum was obtained from all 21 animals at 1 hour, 1 day, and 7 days following the implantation of 22 the antibiotics into the tibiae and serum ampicillin levels were 23 24

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measured using the agar-well diffusion assay described previously
in detail by Bennett et al. A standard curve was constructed
relating the size of the zones of inhibition obtained with a
series of ampicillin standards tested against Sarcina lutea ATCC
series of ampicillin standards tested against oncentrations in the
standard curve.
test sera were then calculated from this standard curve.

RESULTS OF EXAMPLES 1 THROUGH 7

The results of the Immediate Antibiotic Therapy. 7 initial experiment showing the effect of immediate parenteral 8 versus local ampicillin therapy for the prevention of experimental 9 osteomyelitis are presented in Table 2. Radiographic changes were 10 initially detected in the control animals (Groups D, E, and F) at 11 2 weeks postinfection and consisted predominantly of periosteal 12 By 7 weeks, however, the majority of the control 13 animals (75 scores ranging from +5.25 to +7.00 indicating extensive 14 osseous involvement. Radiographic evidence of osteomyelitis was 15 animals that received either a 14 day course of 16 parenteral ampicillin therapy (Group A) or those that received an 17 intramedullary injection of unencapsulated ampicillin (Group C). 18 Only a minimal periosteal reaction was noted at day 42 for Group B 19 injection of intramedullary 20 an microencapsulated ampicillin, however, all other radiographic received 21 parameters were found to be within normal limits. Cultures of the 22 tibiae at 8 weeks following the initiation of treatment showed. 23 that all of the animals treated with either a 14 day course of 24 25

parenteral ampicillin therapy or a single intramedullary injection of microencapsulated ampicillin had sterile bone cultures. Free unencapsulated ampicillin, injected locally into the bone, was ı also effective and sterilized the tibiae of 3 of 4 (75 In contract, 2 all 13 surviving control animals in Groups D, E, and F developed 3 culture-positive osteomyelitis with S. aureus counts ranging from 5 1.3 \times 106 to 2.0 \times 107 CFU recovered per gram of bone. 6 7

Delayed Antibiotic Therapy Without Debridement. Table 3 shows the results of the experiment when antibiotic therapy was delayed for 7 days postinfection and was then initiated without 9 debridement. Of the 8 animals in Group A that received a 14 day 10 course of parenteral ampicillin therapy, 6 (75 <u>aureus</u> bone 11 cultures. Only 2 of these animals survived the entire 12 the experimental protocol; six animals died within 1-2 weeks of 13 completing their antibiotic therapy after developing 14 diarrhea. Of the 7 surviving rabbits in Group C that received an 15 intramedullary application of 45 mg of unencapsulated ampicillin, 16 5 (71 with a single intramedullary application of microencapsulated 17 ampicillin anhydrate (Group B) sterilized the tibiae of 4 of 8 (50 of \underline{s} . aureus recovered from the tibiae of the other animals in 18 19 this group as compared with the controls (Group D). All of the 20 22 control animals developed osteomyelitis with an average of 2.8 x 23 105 CFU of S. aureus recovered per gram of bone. analysis of the proportion of animals in each treatment group with positive bone cultures showed no statistically significant differences among the groups (p = 0.23).

Delayed Antibiotic Therapy With Debridement. In this experiment we evaluated the effect of local antibiotic therapy 2 when used in conjunction with debridement for the treatment of a 7-day established osseous infection. Bacteriological cultures of 3 the tibiae at the time of debridement (before antibiotic therapy 4 was initiated) yielded \underline{S} . aureus in 29 of 30 (97 shown in Table 4, 5 6 all 10 of the animals in Group A that were treated with 7 debridement plus microencapsulated ampicillin sterile bone cultures. In contrast, of the 10 animals in Group B 8 9 that were treated with debridement plus unencapsulated ampicillin 10 7 developed only 3 had sterile bone cultures whereas culture-positive osteomyelitis. A Chi squire analysis showed a 11 statistically significant difference (p < 0.01) in the proportion 12 13 of animals with sterile bone cultures in the microencapsulated ampicillin treated group as compared with the group that was 14 15 treated with the unencapsulated form of the 16 Debridement alone, without local antibiotic therapy, was not 17 effective for the treatment of this established osseous infection 18 with all 10 control animals (Groups C and D) developing 19 20 culturepositive osteomyelitis. 21

Serum Ampicillin Levels. In the experiment where local antibiotic therapy was initiated in conjunction with debridement, serum concentrations of ampicillin were measured for all animals

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application intramedullary microencapsulated ampicillin anhydrate or an equivalent dose of either an 1 unencapsulated free ampicillin. The data is presented in Figure 2 1. Serum levels of ampicillin were only detected at 1-hour after 3 the implantation of the antibiotics into the tibiae. At this time interval, the mean serum concentration of ampicillin in the Group 5 B animals that received 45 mg of unencapsulated ampicillin (0.79 + 6 0.24 micrograms/ml) was approximately 7-fold higher than the mean 7 serum ampicillin concentration of the Group A animals that 8 received an equivalent dose of the microencapsulated form of the 9 10 antibiotic (0.11 + 0.08 micrograms/ml). 11

DISCUSSION RELATED TO EXAMPLES 1 THROUGH 7

Previous attempts to develop a biodegradable antibiotic 12 delivery system for the local treatment of bone infections have 13 met with only limited success. Zilch and Lambiris reported on the 14 treatment of 46 patients with chronic osteomyelitis using a 15 biodegradable fibrin-cefotaxim compound that was implanted into 16 the bone at the time of surgical intervention and reported healing 17 in only 67 limitation of this system was the rapid diffusion of the 18 High concentrations of 19 antibiotic from the fibrin carrier. cefotaxim could only be maintained locally in the would exudate 20 for up to 72 hours. In a separate study, Dahners and Funderburk 21 implanted gentamicin-loaded plaster of paris into the tibiae of 22 rabbits with established staphylococcal osteomyelitis. Although 23 they observed clinical and roentgenographic improvements as 24 25

compared with nontreated controls, nevertheless, 80animals treated developed the gentamicin-loaded plaster of paris 1 culture-positive osteomyelitis. Recently Gerhart et al. evaluated 2 poly(propylenefumarate-co-methylmethacrylate) (PPF-MMA), 3 potential biodegradable carrier for antibiotics. Following the 4 subcutaneous implantation of gentamicin- or vancomycin-loaded 5 cylinders of PPF-MMA in rats, high concentrations of each 6 antibiotic were detected locally in the wound exudate while serum 7 antibiotic levels remained low. Although the PPF-MMA appears 8 promising as a potential biodegradable antibiotic carrier, the 9 efficacy of this system remains to be demonstrated in an 10 experimental animal model of osteomyelitis. 11 12

In the present application we evaluated biodegradable microspheres of poly(DL-lactide-co-glycolide), containing 30.7 13 weight percent ampicillin anhydrate, 14 In the initial osteomyelitis model of the rabbit tibia. 15 experiment where treatment was initiated immediately following the injection of \underline{S} . aureus into the medullary canal, a single 17 intramedullary injection of 100 mg of microencapsulated ampicillin 18 effectively prevented the establishment of osteomyelitis in 100cf Although a 14 day course of the animals tested (Table 2). parenteral ampicillin therapy also prevented osteomyelitis in all animals, the total dose of antibiotic administered to these 22 animals (1,050 mg) was 34 times higher than the dose admimistered 23 to the animals treated locally with the ampicillin-loaded 24 25

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microcapsules (30.7 mg).

In the second experiment, where antibiotic therapy was delayed for 7 days and was instituted without debridement, a 14 2 day course of parenteral ampicillin therapy resulted in a 75 3 treatment failure rate (Table 3). Free unencapsulated ampicillin, 4 implanted locally into the bone, was also ineffective with 71 these 5 animals developing culture-proven osteomyelitis. 6 intramedullary application of microencapsulated ampicillin, on the 7 other hand, sterilized the tibiae of 50 significantly reduced the 8 mean number of S. aureus colonies recovered from the tibiae of the other animals in this group. It is noteworthy that all animals 10 microencapsulated ampicillin anhydrate 11 survived the duration of the experimental protocol without treated locally with 12 In contrast, 6 of 8 (75 developing adverse side-effects. parenteral ampicillin died within 1 to 2 weeks of completing their 14 antibiotic therapy. The cause of death in these animals was most 15 likely antibiotic-induced diarrhea resulting from colonization of 16 the normal intestinal flora by Clostridium difficile, a phenomenon 17 that has been previously noted with rabbits receiving parenteral 18 19 ampicillin therapy. 20

In the final experiment, where local antibiotic therapy
was delayed for 7 days and was instituted in conjunction with
debridement, a 100 animals treated with debridement plus
microencapsulated ampicillin (Table 4). In contrast, of the 10

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animals treated with debridement plus an equivalent dose of unencapsulated ampicillin powder, 70 seen in Figure 5, at 1 hour 1 after implantation of the antibiotics into the medullary canal, 2 the mean serum concentration of ampicillin in the animals 3 receiving unencapsulated ampicillin was approximately 7 times 4 higher (0.79 + .024 micrograms/ml) than in the group that was 5 treated with microencapsulated ampicillin anhydrate (0.11 + 0.08 6 suggests that 7 finding This micrograms/ml). diffuses rapidly from the site of drug unencapsulated administration and does not remain localized for a sufficient time interval to eradicate the infection. The fact that 70 animals 10 treated with the unencapsulated form of the drug developed 11 conclusion. 12 substantiates this ampicillin-loaded microcapsules/spheres, on the other hand, remain osteomyelitis 13 localized at the site of administration thereby continuing to 14 release high concentrations of the antibiotic over time resulting 15 16 in the elimination of the infecting organisms. 17

Applicants' experimental studies have demonstrated that
a controlled-release and biodegradable antibiotic delivery system
was successful for the eradication of a susceptible organism from
an osteomyelitic focus when used in conjunction with adequate
debridement.

Preparation of Ampicillin Anhydrate Microcapsules

23 Preparación SEXAMPLE 8

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About 500 g of a 10 wt alcohol) (PVA) was added to a 1-L (liter) resin kettle and cooled to 5°C while being stirred at 650 1 rpm with a 2.5-in. Teflon turbine impeller driven by a motor and 2 consisting of 5 g of 68:32 3 a control unit. A solution mixture of 40 g of poly(DL-lactide-co-glycolide) in a dichloromethane and 20 g of acetone was prepared in a separate 5 container and stirred magnetically while in an ice bath. In still 6 another container, 5 g of ampicillin anhydrate was dispersed in 15 7 8 This mixture was stirred sonicated to achieve uniform dispersion of single ampicillin g acetone. 9 anhydrate crystals. After sonication, the container was placed in 10 an ice bath, magnetic stirring was continued, and additional 11 acetone was added to give a total of 30 g of acetone. After 12 complete dissolution of the copolymer, the ampicillin-acetone 13 dispersion was added to the copolymer solution. 14 mixture was stirred magnetically in an ice bath for 15 minutes or until homogeneous, and it was then added to the 16 reaction flask containing the aqueous PVA solution. The stir rate 17 was reduced from 650 to 500 rpm after the addition was complete. 18 After 15 minutes, the pressure was reduced to 550 torr to begin 19 slow evaporation of the organic solvent (dichloromethame 20 The pressure was further reduced to 250 torr. This 21 pressure was maintained for another 18 to 24 hours. The flask was 22 then opened, the suspension was removed, and the microcapsules 23 were separated from the PVA solution by centrifugation. 24 microcapsules were then washed twice with water, centrifuged, and 25 26

washed once more with water and recovered by filtration. microcapsules were then dried in vacuo and separated into various 1 size fractions by sieving. A free-flowing powder of spherical 2 3 particles was obtained. 4

EXAMPLE 9

Dissolve 1.2 g of 50:50 poly(DL-lactide-co-glycolide) in 5 102 g of methylene chloride. Ampicillin anhydrate (0.8 g) is next 6 This mixture added to the stirring copolymer solution. 7 (dispersion of drug in the copolymer solution) is then placed in a 8 200-mL resin kettle equipped with a true bore stirrer having a 9 1.5-inch Teflon turbine impeller driven by a motor. 10 mixture is stirring at 700 to 800 rpm, 48 mL of 100 centastoke 11 (cSt) silicone oil is pumped into the resin kettle to cause the 12 poly(DL-lactide-co-glycolide) to coacervate and coat the dispersed 13 ampicillin anhydrate particles. After the silicone oil is added 14 to the resin kettle, the contents of the kettle are poured into 15 heptane to harden the microcapsules/spheres. After stirring in 16 the heptane for 2 hours, the microcapsules/spheres are collected 17 A free-flowing powder of spherical 18 on a funnel an dried. 19 different sized particles is obtained.

In Vitro Characterization of Microcapsules/spheres

The core loadings of microcapsules/spheres comprising 21 ["C]-ampicillin anhydrate and DL-PLG were measured by lignid 22 scintillation counting. The core loading of microcapsules/spheres 23

consisting of unlabeled ampicillin anhydrate and some radiolabeled ampicillin anhydrate and DL-PLG was measured by using a microbial 1 In the former instance, microcapsules/spheres (about 15 2 mg) were solubilized in 1 mL of 0.5 N dimethyl dialkyl quarternary 3 ammonium hydroxide in toluene (Soluene-350) at 55°C for 2 to 4 4 Then, 14 ml of scintillation cocktail (1,4-bis[2-(5-5 phenyloxazolyl] benzene (PPO/POPOP) in toluene) was added, and the 6 scintillation liquid 7 with a radioactivity was measured In the latter instance, microcapsules/spheres 8 spectrometer. (about 15 mg) were placed in 5 mL of methylene chloride. 9 Following dissolution of the DL-PLG excipient, the insoluble 10 ampicillin anhydrate was extracted from the methylene chloride 11 with four volumes of sterile 0.1 M potassium phosphate buffer (pH 12 8.0). These aqueous extracts were then assayed for the antibiotic 13 using Sarcina lutea ATCC 9341 (American Tye Culture Collection, 14 Rockville, MD) and the agar-diffusion microbial assay previously 15 described in the literature by Kavanagh, F. (ed.) Antibiotic 16 Substances in Analytical Microbiology, Vol. II, 1972. 17 18

The in vitro release kinetics of ["C]-ampicillin 20 anhydrate microcapsules/spheres was determined following the 21 placement of 30 mg of microcapsules in an 8-ounce bottle. The 22 release study was initiated by the addition of 50 mL of receiving 23 fluid consisting of 0.1 m potassium phosphate buffer (pH 7.4). 24 The bottle was then sealed and placed in an oscillating (125 cycles/ minutes) shaker bath maintained at 37°C. Periodically, a

- 3-ml aliquot of the receiving fluid was removed for assay and replaced with a fresh 3-ml aliquot of receiving fluid to maintain a constant volume of receiving fluid throughout the study. 3-ml aliquots were assayed for drug by liquid scintillation 2 counting using 12 ml Scinti Verse-I (Fisher Scientific Co., 3 Pittsburgh, PA). The cumulative amount of the drug released into 4 5 6
 - the receiving fluid was calculated. The in vitro release kinetics of unlabeled ampicillin anhydrate microcapsules/spheres was determined in the following 7 8 9 manner: 10
 - anhydrate ampicillin of microcapsules/spheres (about 4 mg of microencapsulated ampicillin anhydrate) and 5.0 ml of sterile receiving fluid (0.1 M potassium 11 phosphate buffer, pH 7.4) were added into dialysis tubing. The 12 ends of the tubing were sealed with plastic clamps. The clamped 13 dialysis tubing containing the microcapsules/spheres were placed 14 into a sterile 8-ounce bottle containing 100 ml of sterile receiving fluid (0.1 M potassium phosphate buffer, pH 7.4). The 19 bottle was placed in a shaker bath maintained at 37°C and shaked 17 20 at 120 cycles per second with about 3-cm stroke. The receiving 21 fluid was previously sterilized in an autoclave for 20 minutes at Several dialysis tubing assemblies were prepared for one 23 release study. At Days 1, 2, 4, 7, 10, 13, 15, 18, and 25, one 24 assembly was removed from its receiving fluid and air dried.

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After drying the assembly, all particles remaining inside the dialysis tubing were quantitatively transferred to a 1 sterile, glass test tube (16 by 125 mm), 5 ml of methylene 2 chloride were added to dissolve the microcapsules, and the drug 3 was extracted with three 5-ml portions of sterile 0.1 M potassium phosphate buffer (pH 8.1). The extraction and preparation of the sample (along with controls) was performed using the procedures 6 for core-loading analysis as discussed above in the extracted 7 samples, and controls using the microbiological assay. Knowing the 8 amount of microencapsulated drug initially placed in the dialysis 9 tubing and the amount of drug remaining in the dialysis tubing 10 after incubation with receiving fluid, the amount of drug released 11 was determined by calculating the difference between them. 12 13

In Vivo Release Profiles of Ampicillin from Microcapsules/spheres The rate and duration of release of ampicillin anhydrate 1 from the microcapsules/spheres were determined in vivo in rats. 2 In one experiment, about 50- to 80-mg doses of microencapsulated 3 and unencapsulated ampicillin anhydrate were sterilized in 4 disposable syringes with a 2.0- or 2.5-Mrad dose of gamma 5 sterile 6 temperature. dry-ice microcapsules/spheres and unencapsulated [14C]-ampicillin anhydrate 7 were then suspended in about 2.0 mL of an injection 8 comprising 2 wt percent of commercially available carboxymethyl 9 cellulose (Type 7LF, Hercules Inc., Wilmington, DE) 10 percent Tween 20 (ICI Americas Inc., Wilmington, DE) in sterile 11 12 15 for 121°C autoclaved at microcapsules/spheres were administered subcutaneously into the 13 14 anesthestized lightly region of Sprague-Dawley rats. The rats were fed standard laboratory food 15 and water ad libidum and were housed in individual stainless steel 16 cages fitted with metabolism funnels and screens that separated 17 and collected the feces and urine. The urine from each rat was 18 collected, weighed, and analyzed for ["C]-content by liquid 19 scintillation counting. The actual doses of microcapsules/spheres 20 or unencapsulated drug administered to each rat was determined 21 after injection by measuring the amount of drug residue in each 22 syringe by liquid scintillation counting. 23 radioactivity excreted daily by each rat was normalized by the 24 dose of microencapsulated or unencapsulated ampicillin anhydrate

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that each rat actually received. This result was then plotted as 1 a function of time. 2

In a second experiment, unlabelled ampicillin anhydrate The rats were microcapsules/spheres were tested in rats. 3 administered the microcapsules/spheres in the same manner as that 4 described in the first experiment. The microbiological assay 5 described above was used to determine the amount of ampicillin in 7 the serum of these rats. 8

In Vivo Efficacy Evaluation of Microcapsules/spheres

Experiments to evaluate the efficacy of prototype 9 microcapsules/spheres in vivo were performed in 250- to 300-g 10 male, Walter Reed strain, albino rats that were anesthetized with 11 The right hind leg was razor-shaved, 12 sodium pentobarbital. scrubbed with Betadine (The Purdue Frederick Co., Norwalk, CT), 13 and swabbed with 70length and 1 cm deep was made in the thigh 14 muscle and contaminated with 0.2 g of sterile dirt. The muscles 15 were traumatized by uniformly pinching them with tissue forceps, 16 and then the wounds were inoculated with known quantities of 17 Staphylococcus aureus ATCC 6538P and Streptococcus pyogenes ATCC 18 19615. All rats were inoculated on the same day of the experiment 19 with the same quantitated bacterial suspension to insure uniform 20 inoculum in all wounds. The artificially contaminated wounds were treated within 1 hour by layering sterile, pre-weighed amounts of microencapsulated antibiotic directly on the wounds. 23

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groups consisted of animals with wounds that either received no placebo 1 with overlaid microcapsules/spheres, or were treated with locally applied, 2 powdered unencapsulated ampicillin anhydrate. 3 treatment, all wounds were sutured closed with 3-0 black silk. 5

Three groups of 20 rats each were used in an efficacy A382-140-1 evaluate Mmicrocapsules/spheres 6 formulated from 70:30 DL-PLG. In this experiment, a group of experiment 7 with wounds overlaid with 0.5 g of unloaded 8 microcapsules/spheres was substituted for the untreated (no therapy) group evaluated in each succeeding dose-response 10 experiment. In addition, a group of 20 rats treated with 0.5 g of 11 ampicillin anhydrate microcapsules/spheres per wound, and a group 12 of 20 rats treated with 120 mg of locally applied uncapsulated 13 ampicillin anhydrate powder per wound were evaluated. 14 animals from each group were sacrificed at 2, 6, 8, and 14 days 15 and evaluated for the presence of ampicillin in the serum and 16 17 tissue and for the presence of infection. 18

Two dose-response experiments were performed in which Microcapsules/spheres A681-31-1, formulated from 70:30 DL-PLG, and 19 Microcapsules/spheres B213-66-1S, formulated from 53:47 DL-PLG 20 were evaluated. Seven groups of 15 rats each were treated with 21 the doses of microcapsules shown in Table I. Each experiment 22 included an additional group of 15 rats which remained untreated. 23 24

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In dose-response Experiment I, five animals from each group were sacrificed at 2, 7, and 14 days and evaluated for 1 ampicillin levels and number of bacteria present per gram of 2 tissue at each wound site. Serum ampicillin levels were assayed 3 at 2, 4, 7, and 14 days. In dose-response Experiment II, five animals from each group were sacrificed at 7, 14, and 21 days and 5 evaluated for ampicillin levels and number of bacteria present per 6 gram of tissue. Serum ampicillin levels were determined at 2, 7, 7 8 14, and 21 days.

Microcapsules/spheres in a 45 to 106 micron size range made by the phase-separation process were evaluated in these experiments. The ampicillin anhydrate content of the microcapsules/spheres (core loading), batch number, and ampicillin anhydrate equivalent for each dose of microcapsules/spheres are shown in Table 1.

In all experiments, bacterial counts were performed on homogenized, preweighed tissue that had been aseptically removed 16 from the wound sites. Serial dilutions of the homogenized tissue 17 specimens were plated on sheep blood agar. 18 Staphylococcus aureus could be easily differentiated from 19 Streptococcus pyogenes on the basis of colonial morphology. 20 Tissue from varying distances around the wound site and serum 21 removed by cardiac puncture were assayed for antibiotic content. 22 This was accomplished by placing discs saturated with known 23 24

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quantities of serum or tissue homogenates on the surface of Mueller-Hinton agar which had been previously seeded with 1 standardized amounts of Sarcina lutea ATCC 9341. Following 2 incubation at 37°C, inhibition zones were measured. 3 diluted stock solutions containing known quantities of ampicillin 4 anhydrate served as standards. Diameters of the inhibition zones 5 were converted to antibiotic concentrations using standard curves 6 generated by plotting the logarithm of the drug concentration 7 8 against the zone diameters. 9

TEST RESULTS

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Microcapsule/spheres In Vitro Evaluation 10

Ampicillin anhydrate was microencapsulated with DL-PLG 11 DL-PLG is a biocompatible aliphatic polyester that 12 undergoes random, nonenzymatic, hydrolytic scission of the ester 13 linkages under physiological conditions to form lactic acid and 14 glycolic acid. These hydrolysis products are readily metabolized. 15 The purpose of the DL-PLG is to control the release of the 16 ampicillin anhydrate from the antibiotic microcapsule/spheres 17 formulation and to protect the reservoir of ampicillin amhydrate 18 from degradation before it is released from the microcapsules/ 19 Two DL-PLG excipients were used in this study. 20 DL-PLG had a lactide-to-glycolide mole ratio of 70:30 and the 21 other, 53:47. The 53:47 DL-PLG biodegrades faster than the 70:30 22 DL-PLG because of its higher glycolide content. 23

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A phase-separation microencapsulation process afforded 95The than 1 better of yields microcapsules/spheres in microencapsulated ampicillin anhydrated product was a fine, 2 The microcapsules/spheres are relatively 3 free-flowing powder. spherical in shape, but have puckered regions. Although these 4 puckered regions exist, the polymer coating was continuous, and 5 there was no evidence of any fractures or pinholes on the surfaces 6 of the microcapsules. Moreover, the photomicrograph obtained by 7 scanning electron microscopy of ampicillin anhydrate microcapsules 8 did not show any evidence of free unencapsulated ampicillin 9 anhydrate crystals either among the microcapsules or protruding 10 11 through the surface of the microcapsules. 12

The drug content (core loading) of the ampicillin anhydrate microcapsule/sphere formations was measured to assess how much ampicillin anhydrate was incorporated in the microcapsules and to determine the bioactivity of the ampicillin anhydrate after it had been microencapsulated.

In general, the core loading of the 45-to 106 microns

size fraction was similar to the theoretical core loading. The

core loading of a few batches of [14C]-ampicillin anhydrate

microcapsules/spheres was determined by microbial assay as well as

by radioassay. Within experimental error, both assays gave

similar results. This indicates that the ampicillin anhydrate was

not inactivated during the microencapsulation process. Also, the

core loading of ampicillin anhydrate microcapsules/spheres was determined by the microbial assay to determine the effect of 2.5 1 Mrad of gamma radiation on the microencapsulated drug. 2 radiation did not inactivate the drug because the core loading 3 remained the same. For instance, 19.3 spheres with 70:30 DL-PLG 4 assayed as 19.0 irradiation and 11.0 DL-PLG assayed as 11.4 5 irradiated unencapsulated and microencapsulated drug were also 6 Irradiated and layer chromatography. 7 nonirradiated samples chromatographed the same, again indicating 8 that no degradation of the drug was caused by gamma radiation at a 10 dose of 2.5 Mrad. 11

In vitro release measurements were used to identify an ampicillin anhydrate microcapsule/sphere formulation that would 12 release all of its drug at a controlled rate over a period of two 13 The formulation that displayed the desired in vitro 14 release kinetics were microcapsules/spheres with diameters of 45 weeks. 15 to 106 microns consisting of about 10 wt percent ampicillin 16 anhydrate (Bristol Laboratories, Syracuse, NY) and microcapsules/ 17 spheres with diameters of 10 to 100 microns consisting of about 35 18 wt percent ampicillin anhydrate (Wyeth Laboratories, West Chester, 19 PA) and about 65 wt percent 53:47 DL-PLG. Figures 3 and 4 show 20 the in vitro release profiles of two samples of these prototype 21 The microcapsules released a desirable initial 22 burst of drug, representing about 30The remaining drug was then microcapsules. 23 released at a slower controlled rate. 24 25

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The in vitro release profile of sterilized (2.5 Mrad), 1

17.6 compared with the release profiles of sterilized (2.0 Mrad), 2

9.6 and 7.8 DL-PLG (Figure 3). 3

Microcapsule/sphere In Vivo Evaluation

performed with were studies 4 unencapsulated ampicillin anhydrate and the same samples of 5 microcapsules that were tested in vitro, as previously described. 6 As shown in Figures 3 and 4, the unencapsulated drug as well as 7 the microcapsules/spheres showed a fast release of drug during Day 8 1. By Day 4, the amount of ampicillin found in the urine or serum 9 of animals dosed with the unencapsulated drug was below the level 10 11 detection of the assay. microcapsule/sphere formulations maintained an elevated level of 12 drug in the urine or serum for extended periods. Both samples of 13 microcapsules/spheres made with the 53:47 DL-PLG had similar 14 release profiles and released drug for about two weeks. 15 illustrated in Figure 5, the microcapsules/spheres prepared with 16 70:30 DL-PLG released drug for at least four weeks. The results 17 of these pharmacokinetic studies corroborate results of the in vivo 18 The 53:47 microcapsules/spheres 19 release studies described. closely meet the desired target duration of release of two weeks. 20 21

The slow rate of ampicillin release from the 70:30 microcapsules/spheres, as shown in Figure 5, may be undesirable 22 because a low level of ampicillin anhydrate released over a long 23 24

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period may provide favorable conditions for the development of drug-resistant bacterial strains. This slower release of drug 1 could be attributed to the slower biodegradation rate of the 70:30 2 DL-PLG, where the water-soluble ampicillin anhydrate remained 3 trapped inside the hydrophobic DL-PLG excipient until the 4 excipient biodegraded completely. 5 microcapsules/spheres prepared with either the 70:30 or 53:47 6 DL-PLG, one could speculate that the release of drug is due to 7 diffusion of the drug through water-filled pores, pores that 8 enlarge as more and more drug is released and as the DL-PLG 10 bioerodes. 11

However, all ampicillin anhydrate microcapsules/spheres formulated effectively reduced bacterial counts in contaminated 12 The most dramatic observation was the rapid elimination 13 of Streptococcus pyogenes. Streptococcus pyogenes was present in wounds. 14 90 from microcapsule/sphere-treated wounds within 48 hours. All 15 three of the microcapsule/sphere batches evaluated were equally 16 successful in eliminating this organism within two days. At 7 17 days Staphylococcus aureus remained in all treated wounds; 18 however, compared to untreated controls, the bacterial count per 19 gram of tissue decreased by at least 2 \log_{10} between Days 2 and 7. 20 This reduction was not observed in untreated controls. In the 21 efficacy evaluation of microcapsules/spheres A382-140-1, wounds 22 treated with unloaded DL-PLG microcapsules, as well as those 23 treated with topical unencapsulated ampicillin anhydrate, remained . 24 25

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infected at 14 days with > 10 organisms per gram of tissue; whereas, 60 ampicillin anhydrate were sterile. The wounds of the 1 4010³ organisms per gram of tissue. By 14 days, 2 regardless of the dose administered (0.5-0.05 g), all wounds remaining 3 treated with microcapsule/sphere sample A681-31-1 were sterile; 4 untreated wounds remained infected with > 105 5 organisms per gram of tissue. At 14 days, all wounds treated with whereas, all 6 0.15 g of microcapsules/spheres B213-66-1S were sterile, however, $5.7 \times 10^2 \frac{1}{5}$ Staphylococcus aureus per gram of tissue were counted in wounds of one animal treated with a 0.25-g dose of encapsulated ampicillin anhydrate. This failure was attributed to 10 an abscess around a suture on the wound surface. All wounds 11 treated with 0.15 g of microcapsules/spheres (B213-66-1S) were 12 however, in the group treated with a 0.05-g dose of 13 microcapsules/spheres, one wound remained contaminated with 3.6 x 14 104 Staphylococcus aureus per gram of tissue. The untreated 15 evaluated 16 averaged 1.4 x 105 animals, control microcapsule/sphere-treated groups, 17 Staphylococcus aureus per gram of tissue. 18 19

20 Serum levels of drug were dependent upon the ampicillin 21 anhydrate reservoir present inside the microcapsules/spheres (core 22 loading), the dose, and the ampicillin release characteristics. 23 Administration of 0.25 g of Microcapsules/spheres A681-31-1, which 24 contained a 45.25 mg ampicillin reservoir per wound, maintained a 25 serum ampicillin level of 8.0 ± 7.3 microgram/milliliter for up to

4 days post-treatment. A dose twice that amount (90.50 mg ampicillin equivalent) maintained detectable serum ampicillin for 1 up to 7 days post-treatment at a serum ampicillin concentration of 2 15.95 \pm 5.0 microgram/milliliter for the first 4 days. 3 ampicillin was not detected in animals whose wounds were treated 4 with microcapsule/sphere doses containing an ampicillin equivalent 5 Even though serum ampicillin was not 6 detected in any animal at 14 days, the tissue levels at this time of 28.50 mg or less. 7 were above the minimal inhibitory concentrations required to kill 8 9 all infecting organisms in This was true with microencapsulated ampicillin anhydrate. 10 microcapsule/sphere doses as low as 0.05 gram per wound. 11 though serum ampicillin was not detected, microbial bioassay for 12 ampicillin in tissue removed from wounds treated with 0.05 gram of 13 microcapsules/spheres (A681-31-1) contained a 14 ampicillin level of 54, 70, and 21 micrograms/gram of tissue at 2, 15 7, and 14 days, respectively. Because the minimal inhibitory 16 concentrations of ampicillin required to kill 95 of Staphylococcus 17 aureus and 97 pyogenes is 0.5. and 0.05 micrograms/milkiliter, 18 respectively, it is a reasonable assumption that a more than 19 adequate therapeutic amount of drug was present at the wound site 20 throughout the two-week treatment period. 21 In vitro release studies performed on microcapsules/ 22 spheres formulated with 70:30 DL-PLG (A382-140-1 and A681-31-1) 23 showed drug release at an efficacious rate over two weeks, but 24 also at a slower rate for an additional 50 days. The continued 25 26

release of low amounts of antibiotic in wounds after two to three weeks is undesirable because of the potential to provide favorable 1 conditions for the emergence of ampicillin resistant organisms in 2 wounds which might harbor small numbers or bacteria. 3 to reduce or eliminate drug trailing, microcapsules/spheres were reformulated by encapsulating ampicillin anhydrate within the 5 faster biodegrading polymer 53:47, DL-PLG (sample B213-66-1S), in 6 vitro release profiles showed a release of 85 to 92 within two 7 weeks. On the seventh day following treatment of wounds with 0.15 8 gram of Microcapsules/spheres B213-66-1S, a mean (n=5) of 162.5 g 9 of ampicillin per gram of tissue was quantitated. In vitro 10 release studies suggest that this amount drops rapidly in the 11 second week so that by 14 days marginal killing concentrations are 12 present. In vivo analysis of tissue removed from wounds treated 13 15 days previously with 0.25 gram of these microcapsules/spheres 14 contained < 1.9 micrograms/gram of ampicillin per gram. Although 15 <0.22 micrograms/gram of ampicillin was detected in wounds treated 16 with 0.15 gram, it was unusual to detect any ampicillin at 14 days 17 in tissue from wounds treated with 0.05 gram per wound. At 21 18 days post-treatment, ampicillin was not detected in any of the 19 20 wounds. 21

No serum levels of ampicillin were detected in any of
the rats treated with Microcapsules/spheres B213-66-15. This was
expected because lower doses (ampicillin equivalent) were
administered. (Table 1).

B. Cefazolin (CZ) microspheres. The CZ microspheres used in these studies were produced by Southern Research Institute, 1 The microspheres consisted of 77.8 weight } 2 copolymer (50:50 molar ratio of lactide to glycolide) with a core 3 leading dose of 22.2 weight & cefazolin. microspheres ranged from 90 to 355 um in diameter and they were 5 In vitro release sterilized with 2.7 Mrad of gamma radiation. 6 kinetic studies showed that approximately 20% of the cefazolin was 7 released from the microspheres within 6 hours, with the remainder 8 of antibiotic release extending over a period of 15 days. 9 10

Experimental wounds were Rat wound infection model. surgically created in the paraspinous muscles of Sprague-Dawley 11 rats following induction of anesthesia with ketamine and xylazine. 12 Sterile sand (100 mg) was implanted into the wound site to simulate 13 a foreign body and the wounds were inoculated with 5 \times 10 6 CFU each 14 of Staphylococcus aureus ATCC 27660 and Escherichia coli ATCC 15 25922. The minimum inhibitory concentration (MIC) of cefazolin for 16 each of these organisms was 4 ug/ml and 2 ug/ml, respectively. The 17 animals were then randomly distributed in 6 groups. Groups A, B, 18 and C (6 rats per group) received local antibiotic therapy with 50 19 mg, 250 mg, or 500 mg of CZ microspheres, respectively. The 20 microspheres were applied directly to the wounds and care was taken 21 to achieve a relatively uniform distribution of the drug throughout 22 the wound site. Group D (6 rats) received local antibiotic therapy 23 with 110 mg of CZ powder. This dose was equivalent to the core-24 25

loading dose of cefazolin contained in 500 mg of CZ microspheres used to treat the Group C animals. Group E (6 rats) received 1 systemic antibiotic therapy with cefazolin (30 mg/kg) which was 2 administered as a single intramuscular bolus immediately after 3 bacterial contamination of the wounds. Group F (3 rats) served as 4 controls and received no antibiotic therapy. The wounds were then 5 closed with surgical staples and the animals were returned to their 6 cages. On postoperative day # 28, the rats were euthanized and 7 tissue was obtained from each wound for quantitation of surviving 8 bacteria. The tissue was weighed, homogenized, and serial 10-fold 9 dilutions were prepared and plated on blood agar. The number of 10 bacteria recovered from each wound was quantitated and expressed as 11 12 CFFU/g tissue. 13

Rabbit fracture-fixation model. This study was conducted in two segments and was designed to evaluate the effect of early as well 14 as delayed local antibiotic therapy for the prevention of infection 15 in experimental fractures. In segmentI, open fractures were created 16 in the right tibiae of New Zealand White rabbits after induction. 17 of anesthesia with ketamine and xylazine. The fractures were then 18 inoculated with 0.5 ml of S. aureus ATCC 27660 (2.0 \times 10⁷ CFU/ml). 19 Within 30 minutes following bacterial contamination, the animals 20 were randomly distributed in 5 groups. Group A (8 rabbits) received 21 local antibiotic therapy with 300 mg of cefazolin microspheres 22 which was applied directly to the fracture site and the deep 23 musculature. Group B (8 rabbits) received local antibiotic therapy 24 25

with an equivalent dose of CZ powder. Group C (8 rabbits) received systemic antibiotic therapy with cefazolin (25 mg/kg/day) for 7 ı days. Groups D and E (4 rabbits per group) served as controls and 2 received either local application of placebo microspheres (without 3 cefazolin) or no treatment, respectively. The fractures were then 4 reduced and plated with a 4-hole dynamic compression plate. 5 Immediately prior to wound closure, animals in Groups A and B 6 received an additional dose of either CZ microspheres (300 mg) or 7 an equivalent dose of CZ powder, respectively, which was applied 8 directly over the fixation plates and the periosteal tissue. The 9 wounds were then repaired with sutures and the animals were 10 returned to their cages. Blood was obtained within 1 hour and 11 again at 24 hours after treatment from all Group A and B animals 12 for quantitation of serum cefazolin levels which was measured by a 13 microbial inhibition bioassay'. Eight weeks later, all swiving 14 animals were euthanized and the tibiae were harvested for 15 bacteriological analysis, the bones were crushed to small pieces 16 with sterile mortar and pestle and saline was added to make a 17 particulate suspension. Serial dilutions were then prepared and 18 streaked on blood agar for bacterial isolation. The number of S. 19 aureus colonies recovered from each specimen was quantitated and 20 21 expressed as CFU/g of bone. 22

In signet II, fractures were created in the right tibia of 29
rabbits and contaminated with S. aureus as described above. After
a 2 hour delay, the animals were randomly distributed in 3 groups.

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Group A (10 rabbits) received local antibiotic therapy with 600 mg of CZ microspheres. Group B (10 rabbits) received local antibiotic 1 therapy with an equivalent dose of CZ powder. Group C (9 rabbits) 2 served as controls and received no treatment. The fractures were 3 then reduced, plated, and the wounds were closed with sutures. 4 Eight weeks later, the surviving animals were euthanized and the 5 tibiae were harvested and processed for isolation of bacteria as 6 7 described above.

Results

Rat wound infection model. Table 5 shows the effect of local versus systemic cefazolin therapy on the contamination rate in rat 10 soft-tissue wounds at 28 days postinfection. Local antibiotic 11 $_{13}$ therapy with CZ microspheres, in doses ranging from 50 to 500 mg 14 per wound, was highly effective for eliminating both organisms from $_{15}$ the wounds. The maximum effect was achieved in the Group ^canimals $_{16}$ who received the highest dose of CZ microspheres (500 mg) where \underline{E} . coli and S. aureus were eliminated from 100% of the wounds. Even at the lowest dose used (50 mg/wound), 4 of 6 wounds were zendered 19 completely sterile. Local antibiotic therapy with free CE peuder 20 sterilized the wounds in 5 of 6 (83%) animals. 21 systemic administration of cefazolin (30 mg/kg failed to sterilize 22 the wounds in any of the 6 Group E animals tested.

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Rabbit fracture-fixation model. Table 6 shows the results of 1 the clinical and bacteriological findings at 8 weeks in 25 2 surviving rabbits when local or systemic antibiotic therapy with 3 cefazolin was initiated within 30 minutes following bacterial contamination of the fractures. Deep infection, defined as the 5 presence of pus on the fixation plate or in the deep tissues, was 6 noted in 6 of the 7 (86%) control animals in Group D (placebo 7 microspheres) and group E (no treatment). Cultures of the tibiae 8 from all 7 controls were positive for S. aureus. 9 surviving Group animals who received a 1 week course of systemic 10 cefazolin therapy, deep infection was noted in 3 cases and S. 11 aureus was recovered from the bones of 4 of the 5 animals. 12 contrast, no clinical evidence of infection was detected in any of 13 the 7 Group A animals who received an equivalent local dose of free 14 CZ powder. Cultures of the tibiae were sterile in 6 of (86%) Group 15 A and 5 of 6 (83%) Group B animals, respectively. There was a 16 statistically significant difference in the mean log S. aureus 17 counts of the Group A and Group B animals and all other groups by 18 analysis of variance (p < 0.05). The mean log \underline{S} . aureus counts for Group C was also significantly different from all groups with the 20 21 exception of Group E (no treatment).

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Table 7 shows the results of the clinical and bacteriological findings at 8 weeks in 23 surviving rabbits when local antibiotic 1 therapy was delayed for 2 hours following bacterial contamination 2 of the fractures. Clinical evidence of infection was present in 5 3 of 7 (71% control animals in Group C and cultures of the tibiae yielded S. aureus in all 7 cases. Of the 8 animals in Group B who 5 received local antibiotic therapy with Cz powder, deep infection 6 was noted in 4 animals and S. aureus was received in 6 of 8 (75%) cases. In contrast, none of the 8 animals in Group Aa (CZ microspheres) developed clinical infections and cultures of the 9 tibiae were sterile in all cases. One way analysis of variance 10 showed a statistically significant difference in the mean log \underline{S} . 11 aureus counts between Groups A and B (p = 0.0014); Groups A and C 12 (p < 0.0001); and Groups B and C (p = 0.0269). 13 14

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- 1 11. C. Heisterkamp, J. Vernick, R.L. Simmons, and T. Matsumoto,
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- Applicants have developed microencapsulated antibiotics for the local treatment of containinated surgical and traumatic 3 wounds. Preliminary studies have shown that local application of 4 biodegradable antibiotic microspheres to experimental wounds that 5 were contaiminated with resistant bacteria was extremely effective 6 for prevention of wound infection. This success is attributed to 7 the significantly higher local tissue antibiotic levels that can be 8 achieved at the wound site with direct local application of 9 microencapsulated antibiotics as compared to conventional systemic 10 antibiotic dosing. The findings of the experimental studies are 11 12 summarized below: 13
 - 1. Ampicillin microspheres effectively prevented infection in 8/11 (73%) animals whose wounds were inoculated with an ampicillin-resistant strain of s. aureus (MIC = 750 ug/ml). Systemic ampicillin failed in 9/9 (100%) cases.
 - 2. Cefazolin microspheres effectively prevented infection in 5/6 (83%) animals whose wounds were inoculated with a methicillin-resistant strain of S. aureus which was also resistant to cefazolin (MIC = 64 ug/ml). Systemic cefazolin failed in 5/6 (83%) cases.

- 3. It is preferred that a initial release (burst) of the encapsulated antibiotic occur within the first day and the 1 remaining antibiotic be released over the next 2 to 3 weeks. 2 3
- EXPERIMENTAL DESIGN FOR RAT SOFT-TISSUE WOUND INFECTION MODEL 4
- Experimental surgical wounds were created in the paraspinous muscle of anesthetized Sprague Dawley rats, each weighing between 5 450 to 550 grams. The wounds were then containinated with 100 mg 6 of sterile sand as an infection-potentiating agent. The wounds 7 were then inoculated with 5 x 10° CFU of S. aureus ATCC 33593. This is a methicillin-resistant strain of S. aureus which is also resistant to cefazolin (MIC = 64 ug/ml). The animals were then 10 assigned to the following treatment groups: 11 12
 - Group A (n = 6): 500 mg of cefazolin (CZ) microspheres was applied directly to the wounds. This dose contained 110 mg of 13 14 cefazolin equivalent. 15
 - Group B (n = 6): 110 mg of free CZ powder was applied 16 directly to the wounds. 17
 - Group C (n = 6): This group received intramecular injections of CZ (30 mg/kg/day) at 8 hour intervals for 7 18 19 consecutive days. . 20

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Group D (n = 3): This group served as controls and did 1 not receive any antibiotic therapy. 2

The wounds were then closed with surgical staples and the animals were returned to their cages for the next 5 weeks. At that 3 time, the animals were humanely euthanized and tissue was removed 4 from the wounds and cultured for the presence of bacteria. The 5 bacteriological data are presented in Table 8. 7

VIII. UTILITY

Successful controlled release of bioactive ampicillin 8 anhydrate was achieved in vitro and in vivo. 9 microcapsules/spheres effectively controlled or eliminated 10 Staphylococcus aureus and Steptococcus pyogenes from infected 11 wounds in rats. Additionally, the formulation would be effective 12 in the treatment of all bacterial infections caused by organisms 13 sensitive to the antibiotic encapsulated including but not limited 14 Enterobacteriaceae; Klebsiella sp.; Bacteroides sp.; 15 Enterococci; Proteus sp.; Streptococcus sp.; Staphylococcus sp.; 16 Pseudomonas sp.; Neisseria sp.; Pedptostreptococcus sp.; 17 Fusobacterium sp.; Actinomyces sp.; Mycobacterium sp.; Listeria 18 sp.; Corvnebacterium sp.; Proprionibacterium sp.; Actinobacillus 19 sp.; Aerobacter sp.; Borrelia sp.; Campylobacter sp.; Cytophaga 20 sp.; Pasteurella sp.; Clostridium sp.; Enterobacter aerogenes; 21 vulgaris; 22 proteus Staphylococcus aureus; Streptococcus polygenes; Actinomyces sp.; Peptococcus sp.; 23 24

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Campylobacter fetus; and Legionella pneumophila. Results
indicate that optimal microcapsules/spheres should exhibit a
programmed release of an appropriate concentration of antibiotic
over about a 14 day to about a 6 week time period after which
time the microcapsule/sphere should biodegrade, leaving no trace
of drug or excipient.

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PHASE II

This illustrative phase of this invention relates to a novel
pharmaceutical composition, a microcapsule/sphere formulation,
may contain a pharmaceutically-acceptable adjuvant that comprises
an antigen encapsulated within a biodegradable polymeric matrix
such as poly

	(DL-lactide-co-glycolide) (DL-PLG), wherein the relative ratio between the
	(DL-lactide-co-glycolide) (DL-r Lo), lactide and glycolide component of the DL-PLG is within the range of 90:10 to
	lactide and glycolide component of the offertive pretreatment of animals
	lactide and glycolide component of animals 0:100, and its use, as a vaccine, in the effective pretreatment of animals
	0:100, and its use, as a vaccine, or (including humans) to prevent intestinal infections caused by a virus or (including humans) to prevent intestinal infections caused by a virus or
	of this invention, applicants round
	allows E. coli RDEC-1 to attach to the
	that enhances the initial series are initial series and initial series are series
:	intestinal mucosa, applicants in the
	antigens that contact the intestinal in-
	homogeneously dispersing AF/R1 pin water homogeneously dispersing AF/R1 pin water parch localization. New Zealand
•	included a size range selected for Peyer's Patch localization. New Zealand
	white rabbits were primed twice with 50 micrograms of either
	the nonencapsulated AF/RI by endoscope
	sheld rissues were removed and centural pro-
	and synthetic AF/R1 pepudes were in
i	processored possible I and/or b corrections
5	The synthetic peptides represented properties and theoretical criteria. In were selected from the AF/R1 subunit sequence using theoretical criteria. In
7	rabbits which had received nonencapsulated AF/R1, Peyer's Patch cells
8	rabbits which had received nonencapsular in vitro in response to AF/R1
19	demonstrated slight but significant proliferation in vitro in response to AF/R1
20	a rip i cynthetic peptides. In labora water
	A E/D1 Pever's Patch cells demonstrated
21	and the synthetic pepulate.
22	nodes responded similarly
23	spleen and mesenteric lymph houes response to the synthese peptide groups of animals, while there was a greater response to the synthese peptide
24	groups of animals, within

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	microencapsulated AF/R1. These
1	AF/R1 40-55 in rabbits that had received microencapsulated AF/R1. These
2	AF/R1 40-55 in rabbits that had received of AF/R1 potentiates the mucosal data demonstrate that microencapsulation of AF/R1 potentiates the mucosal data demonstrate that microencapsulation of AF/R1 potentiates the mucosal
3	data demonstrate that microencapsulation delivers and its linear peptide cellular immune response to both the native protein and its linear peptide
	antigens.
4	characterized by antipilus
	A primary mucosal immune response, characterized by antipilus
•	a phire with E. coli RDEC-1. However, induction of
7	IgA, follows infection of rations with pilus antigen an optimal primary mucosal response by enteral vaccination with pilus antigen
8	depends on immunogenicity of pilus protein, as well as such factors as its
9	depends on immunogenicity of pilus proteins
10	ability to survive gastrointestinal tract (GI) transit and to target
11	immunoresponsive tissue. We tested the effect of incorporating AF/RI pilus
12	antigen into resorbable microspheres upon its ability to induce primary mucosal
13	a responses after direct inoculation and
	i-soculated with 50 micrograms
14	a miled into uniformally sized (5 70 miles)
15	antigen alone of incorporation was by
16	microspheres (MIC) of poly(DL-lactide-coglycolide). Inoculation was by
17	intra-duodenal (ID) intubation via endoscopy or directly into the ileum near a
18	intra-duodenal (ID) intubation via a procedure (with the cecum ligated to enhance Peyer's patch via the RITARD procedure (with the cecum ligated to enhance).
	and a reversible iteal the to store
19	send at 2 weeks for collection of get
20	bled and nurged weekly for 5 weekly
21	obtain gut secretions. Anti-pilus IgA and IgG were measured by ELISA.
22	obtain gut secretions. Anti-pilus 15.5
23	TABLE9
24	RESULTS: *pos/test RITARD-PILI RITARD-MIC ID-PILI ID-MIC.

	(m. !.4\) #*	7/8	4/8	1/2	0/3
1	Anti-pilus IgA (Ilulu)	0/8	. 3/8	0/2	1/3
2	Anti-pilus IgG (serum) Native pilus antigen l			ine in 7/8 R	ITARD
3	Native pilus antigen l	ed to a 1	100321 167- 2012	the groups	were not
4	rabbits. MIC caused a similar	r respon	se in only 4/6, but	weemic Toli	i responses
5	statistically different. MIC (t	ut not p	ili) induced some s	systeme 260	- couleted
6	(highest in animals without m	ucosal r	esponses). Results	in rabbits i	nocuiziei
	- similar for pili, but	no muco	osal response to ID	-MIC was in	oics.
7	THE STARY Inoculation wi	ith pilus	antigen produces a	primary mi	ncozat 18W
8	response. Microencapsulation	n does n	ot enhance this res	ponse, altho	ugh the
9	antigen remains immunogen	ic as sho	own by measurable	mucosal and	d some
10	antigen remains immunogens strong serum responses. It	must be	determined whether	r priming w	ith antigen
11	strong serum responses. It	musi oc	recognies		
12	in microspheres can enhance	e secono	iary responses.		
13	E		EPITOPE DATA		
14			als and Methods		·
	CFA/I PURIFICA	TION-	INTACT CFA/I p	ili were pun	ined from
15	on con congrue as descri	bed by 1	Hall et al, (1989) [20]. Briefly	, bactena
16	grown on colonization fact	or antig	en agar were subje	cted to shear	ing, with the
17	shearate subjected to diffe	rential C	entrifugation and is	opycnic ban	ding on
18	shearate subjected to difference cesium chloride in the pre-		N-lauryl sarkosine	. CFA/I w	ere dissociated
19	cesium chloride in the pre	Sence U	uci 02 M a	mmonium bi	carbonate (2
20	to free subunits in 6M gu	anididini	lum nci, v.2	(Amicon XI	A 50 simed
21	hr, 25%, passed through a	in ultrafi	iltration memorane	where to I	PRS ON 2 YM
22	hr, 25°), passed through the cell, Danvers, MA), with	concen	tration and buffer e	xchange to	
23	10 stirred cell (Amicon).	Exami	nation of dissociate	d pill by eld	Cuvii
	microscopy demonstrates	i a lack	of pilus structure.		•
24	Illicioscob) and				

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Protein Sequencing-The primary structure of CFA/I has been determined by protein sequencing techniques (Klemm, 1982) and through molecular cloning methods (Karjalainen, et al 1989) [21]. In these two studies there was agreement in all but two of the 147 amino acid residues (at positions 53 and 74). To resolve the apparent discrepancies, CFA/I was enzymatically digested in order to obtain internal amino acid sequence. Trypsin or S. aureus V8 protease (sequencing grade, Boehringer Mannheim) was incubated with CFA/I at a 1:50 w:w ratio (Tris 50 mM, 0.1% SDS, pH 8.5 for 16h at 37° (trypsin) or 24°C (V8)). Digested material was loaded onto precast 16% tricine SDS-PAGE gels (Schagger and von Jagow, 1987) (Novex, Encinitis, CA) and run following manufacturers instructions. Separated samples were electrophoretically transferred to PVDF membranes (Westrans, Schleicher and Schuell, Keene, NH) following Matsiduria (1987) using the Novex minibles apparatus. Blotted proteins were stained with Rapid Coomassie stain (Diversified Biotech, Newton Centre, MA). To obtain the desired fragment containing the residue of interest within a region accessible by automated gas phase sequencing techniques, molecular weights were estimated from standards of molecular weights 20,400 to 2,512 (trypsin inhibitior, myoglobin, and myoglobin cyanogen bromide fragments; Diversified Biotech) using the corrected molecular weights for the myoglobin fragments as given in Kratzin et al., (1989) [22]. The estimated molecular weights for the unknown CFA/I fragments were compared to calculated molecular weights of fragments as predicted for CFA/I from the sequence of CFA/I as analysed by the PEPTIDESORT program of a package developed by the University of

Wisconsin Genetics Computer Group. Selected fragments were cut from the 1 PVDF emebrane and subjected to gas phase sequencing (Applied Biosystem 2 470, Foster City, CA). 3 Monkey Immunization-Three rhesus monkeys (Macaca mulatta) were 4 injected intramuscularly with 250 ug of dissociated CFA/I in complete 5 Freund's adjuvent and subsequently with two injections of 250 ug of antgen in 6 incomplete Freund's adjuvent at weekly intervals. Blood was drawn three 7 weeks after primary immunization. 8 Peptide Synthesis- Continuous overlapping octapeptides spanning the 9 entire sequence CFA/I were synthesized onto polyethylene pins by the method 10 of Geysen et al. [16], also known as the PEPSCAN procedure. Derivitized 11 pins and software were purchased from Cambridge Research Biochemicals 12 (Valley Stream, NY). Fmoc-amino acid pentafluorophenyl esters were 13 purchased from Peninsular Laboratories (Belmont, CA), 14 1-hydroxybenzotriazole monohydrate (HYBT) was purchased from Aldrich, and reagent grade solvents from Fisher. To span the entire sequence of CFA/I 15 16 with a single amino acid overlap of from one peptide to the next, 140 total pins were necessary, with a second complete set of 140 pins synthesized 17 18 simultaneously. 19 ELISA procedure- Sera raised in monkeys to purified dissociated pik 20 were incubated with the pins in the capture ELISA assay of Geysen et al., [16] 21 with the preimmune sera of the same animal tested at the same dilution simultaneously with the duplicate set of pins. Dilution of sera used on the pins 22 23

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was chosen by initial titration of sera by standard ELISA assay and immunodot blot assay against the same antigen.

RESULTS

It was essential to utilize the correct sequence of CFA/I in the synthesis of the pins for both T- and B-cell experiments to carry out the studies as planned. At issue were the amino acids at position 53 and 74; incorrect residues at those positions would effect 36 of 138 pins (26%) for T-cell epitope analysis and 30 of 140 pins (21%) for B-cell analysis. To resolve the discrepancy in the literature, purified CFA/I was proteolytically digested separately with trypsin and with S. aureus V8 protease (V8). These enzymes were chosen in order to give fragments with the residues of interest (53 and 74) relatively near to the N-terminus for automated Edman degradation (preferably 1-15 residues). These digests were separated on tricine SDS-PAGE gels (Fig 24 A) and molecular masses of fragments estimated. A fragment of 3459 calculated molecular mass is expected from the trypsin digest (corresponding to amino acids 62-94) and a fragment of 5889 calculated molecular mass is expected from the V8 digest (residues 42-95). These fragments were located within each digest (arrows in Fig.24), and a companion gel with four lanes of each digest was run, electrophoretically transferred to PVDF, the bands excised and sequenced. N-terminal sequences of each fragment are given in Fig.24 B. The N-terminal eighteen residues from the trypsin fragment were determined that corresponded to positions. 62-79 in CFA/I. Position 74, a serine residue was consistent with that determined by Karjalainen et al., (Karjalainen et al., 1989). Nineteen midus

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of the V8 fragment were determined, corresponding to residues 41-60 of the parent protein. The twelfth residue of the fragment contained an aspartic acid, also consistent with Karjalainen et al., (1989). All other residues sequenced were consistent with those published previously (including residues 1-29, not shown). For the following peptide synthesis were therefore utilized the complete amino acid sequence of CFA/I consistent with Karjalainen et al., (1989).

Sera from monkeys immunized with CFA/I subunits were tested in a modified ELISA assay, with the preimmunization sera tested simultaneously with duplicate pins. Assays results are displayed in Fig. 25. Monkey 2Z2 (fig. 2A) responded strongly to six regions of the CFA/I sequence. Peptide 14 (the octapeptide 14-21) gave the strongest response with four pins adjacent to it (11, 12, 13, and 15) also appearing to bind significant antibody. The other 2Z2 epitopes are centered at peptides 3, 22, 33, 93, and 124. Monkey 184D (Fig. 17B) also responded strongly to peptide 14, although the maximum response was to peptide 13, with strong involvement of peptide 12 in the epitope. Additional epitopes recognized by 184d were centered at peptides 22, 33, 66, and 93. The third monkey serum tested, 34, responded to this region of the CFA/I primary structure, both at peptides 1, 12 and weakly at 14. Two other epitopes were identified by 34, centered at peptides 67 and 128. Figure 26 illustrates the amino acids corresponding to the epitopes of CFA/I as defined by the response of these three monkeys aligned with the entire primary structure. The entire antigenic determinants are mapped and areas of overlap

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criteria published by Rothbard and Taylor [7]. The sequence numbers of the first amino acid of the predicted segments are shown in Table 1.

Lymphocyte proliferation of monkey spleen cells to CFA/I synthetic peptides. To determine which segments of the CFA/I protein are able to stimulate proliferation of CFA/I immune primate lymphocytes in vitro, three Rhesus monkeys were immunized with CFA/I subunits, and their splenic lymphocytes were cultured with synthetic overlapping decapeptides which represented the entire CF/I sequence. Concentrations of peptides used as antigen were 6.0, 0.6, and 0.6 ug/ml. Proliferative responses to the decapeptides were observed in each of the three monkeys (fig.9-12. The majority of the responses occurred at the 0.6 and 0.06 ug/ml concentrations of antigen and within distinct regions of the protein (peptides beginning with residues 8-40, 70-80, and 27-137). A comparison of the responses at the 6.0, 0.6 and 0.06 ug/ml concentrations antigenic peptide for one monkey (2&2) are shown (fig.12-14. Taking into account all concentrations of antigen tested, spleen cells from monkey 184D demonstrated a statistically significant response to decapeptides beginning with CFA/I amino acid residues 3, 4, 8, 12, 15, 21, 26, 28, 33, 88, 102, 10, 133, 134, and 136 (fig. 27. Monkey 34 had a significant response to decapeptides beginning with residues 24, 31, 40, 48, 71, 72, 77, 78, 80, 87, and 102, 126 and 133 (Fig.28); monkey 222 responded to decapeptides which began with residues 4, 9, 11, 12, 13, 14, 15, 16, 17, 20, 27, 35, 73, 79, 18, 127, 129, 132, and 133 (fig.27). Peptides beginning with amino acid residues 3 through 2 were synthesized with either a glutamic acid or an asparagine substituted for the aspartic acid residue at

position twelve to prevent truncated peptides. The observed responses to peptides beginning with residue 8 (monkey 184d), and residues 9, 11, 12 (monkey 2Z2) occurred in response to peptides that had the glutamic acid substitution. However, the observed responses to peptides beginning with residue 3, 4, and 12 (monkey 184D), a well as residue 4 (monkey 2Z2) occurred in response to peptides that had the asparagine substitution. Monkey 34 did not respond to any of the peptides that had the substitution at position twelve. All other responses shown were to the natural amino acid sequence of the CFA/I protein. Statistical significance was determined by comparing the cpm of quadruplicate wells cultured with the CFA/I peptides to the cpm of wells cultured with the CFA/I peptides.

Analysis of decapeptides that supported proliferation of lymphocytes from CFA/I immune animals. Of the 39 different peptides that supported proliferative responses, thirty contained a serine residue, 19 contained a serine at either position 2, 3, or 4, and nine had a serine specifically at position 3. Some of the most robust responses were to the peptides that contained a serine residue at the third position. The amino acid sequence of four such peptides is shown in Table 3.

VII. DETAILED DESCRIPTION OF THE INVENTION

Applicants have discovered efficacious pharmaceutical compositions wherein the relative amounts of antigen to the polymeric matrix are within the ranges of 0.1 to 1.5% antigen (core loading) and 99.9 to 98.5% polymer, respectively. It is preferred that the relative ratio between the lactide and

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glycolide component of the poly(DL-lactide-co-glycolide) (DL-PLG) is within the range of 90:10 to 0:100. However, it is understood that effective core loads for certain antigens will be influenced by its microscopic form (i.e. bacteria, protozoa, viruses or fungi) and type of infection being prevented. From a biological perspective, the DL-PLG or glycolide monomer excipient are well suited for in vitro drug (antigen) release because they elicit a minimal inflamatory response, are biologically compatible, and degrades under physiologic conditions to products that are nontoxic and readily metabolized. Surprisingly, applicants have discovered an extremely effective method for the protection against bacterial or viral infections in the tissue of a mammal (human or nonhuman animal) caused by enteropathogenic organisms comprising administering orally to said animal an immunogenic amount of a pharmaceutical composition consisting essentially of an antigen encapsulated within a biodegradable polymeric matrix. When the polymeric matrix is DL-PLG, the most preferred relative ratio between the lactide and glycolide component is within the range of 48:52 to52:48. The bacterial infection can be caused by bacteria (including any derivative thereof) which include Salmonella typhi, Shigella sonnei, Shigella flexneri, Shigella dysenterise, Shigella boydii, Escheria coli, Vibro cholera, yersinia, staphylococcus clostridium and campylobacter. Representative viruses contemplated within the scope of this invention, susceptible to treatment with the above-described pharmaceutical compositions, are quite extensive. For purposes of illustration, a partial listing of these viruses (including any derivative thereof) include hepatitis A, hepatitus B, rotaviruses, polio virus human immunodeficiency

	viruses (HIV), Herpes Simplex virus type 1 (cold sores), Herpes Simplex virus
1	Varicella-zoster virus (chicaci pour
2	type 2 (Herpesvirus gemans), Epstein-Barr virus (infectious mononucleosis; glandular fever; and Burkittis
3	
4	lymphoma), and cytomegalo viruses. A further representation description of the instant invention is as
5	A further representation description
6	follows: A. (1) To homogeneously disperse antigens of enteropathic
7	A. (1) To homogeneously early Organisms within the polymeric matrix of biocompatible and biodegradable organisms within the polymeric matrix of biocompatible and biodegradable
8	organisms within the polymeric matrix or microspheres, 1 nanogram (ng) to 12 micross in diameter, utilizing equal
9	microspheres, 1 nanogram (fig) to 12 microsphere
10	molar parts of polymerized faction and so within the range of about 0.1 to 52:48 DL-PLG) such that the core load is within the range of about 0.1 to
11	52:48 DL-PLG) such that the object of the dispersed antigen can then 1.5% by volume. The microspheres containing the dispersed antigen can then
12	be used to immunize the intestine to produce a humoral immune response
13	composed of secretory antibody, serum antibody and a cellular immune response is
14	response consisting of specific T-cells and B-cells. The immune response is
15	directed against the dispered antigen and will give protective immunity against
16	directed against the dispered antigen and with general derived.
17	the pathogenic organism from which the antigen was derived.
18	the pathogenic organism to the pathogenic organi
	coli RDEC-1 to attach to rabbit intestinal brush borders thus promoting
19	AF/R1 pilus protein was nome
20	matrix of biocompatible and biocogn
21	de lieune in diameter (Figure 9 and photograph)
22	microspheres, 1-12 micross in calculations and glycolide (50:50 DL-PLG) such equal molar parts of polymerized lactide and glycolide (50:50 DL-PLG) such
23	equal molar parts of pulymentation
24	that the core load was .62% by weight.

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	(3) The microspheres were found to contain immunogenic
1	- Lhit spleen (Figure 11) and Peyer's parts (1.1821)
2 .	AF/R1 by immunizing both fabble specific IgM
3	AF/R1 by immunizing both rabbit specific IgM 3) B-cells in vitro. The resultant cell supernatants contained specific IgM The antibody response was comparable
4	antibody which recognized the AF/RI. The analysis
5	containing 30 interest
6	(4) Microspheres conditions of two separate used to intraintestinally (intraduodenally) immunize rabbits on two separate
7	one week later, compared
8	ta tiente pever s patente
9	response to AF/RI will
10	-0 04 and 110-14-2 2)
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12	transformation (F-cells) (Figures 2 and 15) (Figures 14 and 15). Similarly enhanced B-cell responses were also detected in
13	(Figures 14 and 15. Similarly ennanced 2 - Cell response was also detected
14	(Figures 14 and 15. Similarly emiantees to one AF/R1 peptide fragment,
	and the Spice it to one
15	The cellular initiation of
16	er recretory altitudes
17	ancelory allucos
18	Table 1); but indicates that a secretary that the rabbits so immunized could be protected upon challenge with the E
19	
20	Coli RDEC-1. B. Microspheres do not have to be made up just prior to use as
21	B. Microspheres to the not been effective in rabbits for
22	B. Microspheres do not been effective in rabbits for with liposomes. Also liposomes have not been effective in rabbits for
23	with liposomes. Also of lipopolysaccharide antigens. intestinal immunization of lipopolysaccharide antigens.

1	C. (1) Only a small amount of antigen is required (ugs) when
2	dispersed within microspheres compared to larger amounts (mgms) when
	antigen is used alone for intestinal immunization.
3	(2) Antigen dispersed within microspheres can be used orally
4	for intestinal immunization whereas antigen alone used orally even with gastric
5	acid neutralization requires a large amount of antigen and may not be effective
6	
7	for intestinal immunization.
8	(3) Synthetic peptides with and without attached synthetic
9	adjuvants representing peptide fragments of protein antigens can also be
10	dispersed within microspheres for oral-intestinal immunization. Free peptides
11	would be destroyed by digestive processes at the level of the stomach and
12	intestine. Any surviving peptide would probably not be taken up by the
13	intestine and therefore be ineffective for intestinal immunization.
14	(4) Microspheres containing antigen maybe placed into
15	gelatin-like capsules for oral administration and intestinal release for improved
16	intestinal immunization.
17	(5) Microspheres promote antigen uptake from the intestine and
18	the development of cellular immune (T-cell and B-Cell) responses to antigen
19	components such as linear peptide fragments of protein antigens.
20	(6) The development of intestinal T-cell responses to antigens
21	dispersed within microspheres indicate that T-cell immunological memory will
22	be established leading to long-lived intestinal immunity. This long-lived
23	intestinal immunity (T-cell) is very difficult to establish by previous means of
24	intestinal immunization. Failure to establish long-lived intestinal immunity is a

	fundamental difficulty for intestinal immunization with non-viable antigens.
1	and the sectional long-lived immunity only a short lived secretory minorary
2	response is established lasting a few weeks after which no significant
3	response is established lasting a lew weekling
4	immunological protection may remain.
5	D. (1) Oral intestinal immunization of rabbits against E. coli
6	RDEC-1 infection using either whole killed organisms, pilus protein
	i as as linopolysaccharide preparations.
7	(2) Microspheres containing adherence pilus protein ra rich
8	its antigen peptides for oral intestinal immunization of rabbits against RDEC-1
9	its antigen peptides for that interest antigen peptides for that interest and its antigen peptides for the interest and its antigen period in the interest and its antig
10	infection.
11	(3) Oral-intestinal immunization of humans against
	enterotoxigenic E. colj infection using either whole killed organisms, pilus
12	in accountions or lipopolysaccharide preparations.
13	protein preparations of april proteins of april proteins CFA/I, (4) Microspheres containing adherence pilus proteins CFA/I,
14	(4) Microspheres on a intestinal immunization of
15	II, III and IV or their antigen peptides for oral intestinal immunization of
16	humans against human enterotoxigenic E. coli infections.
	(5) Oral-intestinal immunization of humans against other
17	enteric pathogens as salmonella, shigella, camphlobacter, hepatitis-A virus,
18	
19	rota virus and polio virus. (6) Oral-intestinal immunization of animals and humans for
20	(6) Oral-intestinal immunization of annual tree
	mucosal immunological protection at distal mucosal sites as the bronchial tree
21	in lungs, genito-urinary tract and breast tissue.
22	IU inugo, Reinto

	E. (1) The biocompatible, biodegradable co-polymer has a long
1	E. (1) 210
2	history of being safe for use in humans since it is the same one used in
3	resorbable suture material. (2) By using the microspheres, we are now able to immunize
	(2) By using the microspheres, we are improved for
4	(2) By using the interest. the intestine of animals and man with antigens not normally immunogenic for
5	shey are either destroyed
6	the intestinal mucosa because they are or only weakly immunogenic if taken
7	the intestinal mucosa because they are only weakly immunogenic if taken to be taken up by the intestinal mucosa or only weakly immunogenic if taken
8	up. (3) Establishing long-lived immunological memory in the
9	(3) Establishing to a sering immunized using microspheres.
10 -	intestine is now possible because T-cells are immunized using microspheres. (4) Antigens that can be dispersed into microspheres for
11	(4) Anugens mar can proteins, glycoporteins,
12	intestinal immunization include the following: proteins, glycoporteins,
	synthetic polysactimates
13	C PS) synthetic lipopolysacting section of the control of the cont
14	annihetic muramyi dipepuda 2
15	attached adjuvants such as synthesis that the attached adjuvants such as some attached adjuvants and the attached adjuvants such as some attached adjuvants and the atta
16	(5) The subsequent minimum Pever's patch) by the
	systemic (spleen and serum antibody) or local (intestine, Peyer's patch) by the
17	for the intestinal intestinal interestinal i
18	main within macrophage course
19	5-10 microns in diameter remain was a local intestinal immune response.
20	5-10 microns in diameter remain was local intestinal immune response. Peyer's patch in the intestine and lead to a local intestinal immune response.
	s —icrons in diameter leave die s
21	and migrate to the mescular sympanic
22	within macrophages and impor- spleen resulting in a systemic (serum antibody) immune response.
23	spleen resulting in a system v

	(6) Local or systemic antibody mediated adverse reactions
1	because of preexisting antibody especially cytophyllic or IgE antibody may be
2	because of preexisting antibody especially because of their being minimized or eliminated by using microspheres because of their being
3	minimized or eliminated by using interosphere
4	phagocytized by macrophages and the antigen is only available as being phagocytized by macrophages and the antigen is only available as being attached to the cell surface and not free. Only the free antigen could become
5	attached to the cell surface and not tree. Only
6	attached to the cell surface and to the surface of mast cells resulting in attached to specific IgE antibody bound to the surface of mast cells resulting in
7	attached to specific light animos in attached to specific light animos in a specific light animos in attached to specific
8	anaphylaxis.
9 ,	anaphylaxis. (7) Immunization with microspheres containing antigen leads
10	to primarily IgA and IgG antibody responses rather than an IgE antibody
11	to primarily IgA and IgO allows and IgO allows are response, thus preventing subsequent adverse IgE antibody reactions upon
12	ALA ARTIGER
	the above, the encapsulation of the following symmetry
13	peptides are contemplated and considered to be well within the scope of this
14	
15	invention:
16	(1) AF/R1 40-55;
17	(2) AF/R1 79-94;
18	(3) AF/R1 108-123;
19	(4) AF/R1 1-13;
20	(5) AF/R1 pepscan 16AA;
21	(6) CFA/I 1-13; and
22	(7) CFA/I pepscan 16AA.
23	(8) Synthetic Pepetides Containing CFA/I Pilus Protein
24	T-cell Epitopes (Starting Sequence # given)

	
1	4(Asn-Ile-Thr-Val-Thr-Ala-Ser-Val-Asp-Pro),
2	8(Thr-Ala-Ser-Val-Asp-Pro-Val-Ile-Asp-Leu),
3	12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gin-Ala-Asp),
4	15(Ile-Asp-Leu-Leu-Gln-Ala-Asp-Gly-Asn-Ala),
	20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
5	26(Pro-Ser-Ala-Val-Lys-Leu-Ala-Tyr-Ser-Pro),
6	72(Leu-Asn-Ser-Thr-Val-Gln-Met-Pro-Ile-Ser),
7	78(Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Gln),
8 .	87(Gln-Val-Leu-Ser-Thr-Thr-Ala-Lys-Glu-Phe),
9	126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr), and
10	133(Gly-Asn-Tyr-Ser-Gly-Val-Val-Ser-Leu-Val), and mixtures
11	
12	thereof. (9) Synthetic Peptides Containing CFA/I Pilus Protein B-cell (antibody)
13	
14	Eptiopes (Starting Sequence # given)
15	3(Lys-Ana-Ile-Thr-Val-Thr-Ala-Ser-Val),
16	11(Val-Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
17	22(Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
18	32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-Thr-Phe-
	Glu-Ser-Tyr-Arg-Val),
19	32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe),
20	38(Lys-Thr-Phe-Glu-Ser-Tyr-Arg-Val),
21	66(Pro-Gin-Leu-Thr-Asp-Val-Leu-Asn-Ser),
22	93(Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala),
23	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr),
24	174(F)3. v.m. v.m. v

	127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
1	127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr- 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-
2	124(Lys-Thr-Ala-Gly-Inr-Ala-Tiber)
	Ser), and mixtures thereof.
3	Ser), and mixing CFA/I pilus protein T-cell and (10) synthetic peptides containing CFA/I pilus protein T-cell and
4	enitores (Starting Sequence # given)
5	The Val-The-Ala-Sci 200
6	3(Lys-Asn-He-Thi-Val - 1986), 8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gin-Ala-Asp),
7	8(Inr-Ala-Sc Leu-Bal-Ile-Asp-Leu-Leu-Gin-Ala-Asp), 11(Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gin-Ala-Asp),
8	11(Bal-Asp-Pro-Ball and Pro-Ser-Ala-Val), 20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
9	20(Ala-Asp-Gly-Asn-Ala-Do-Thr-Ala-Gly-Asn-Tyr-Ser), and 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and mixtures
10	124(Lys-Thr-Ala-Gly-Thr-Ala-Flo Cly-Asn-Tyr-Ser), and mixtures
11	124(Lys-1 nr-Ala-Oi) 2400 124(Lys-1 nr-Ala-O
	thereof.
12	
	(11) synthetic peptides containing CFA/I pilus protein T-cell and
13	B-cell (antibody) epitopes (Starting Sequence # given)
14	B-cell (anubody) epitopes (
	CFA/I pilus protein T-cell epitopes
15	CFA/L DIVIS DE LA COMPANIA DEL COMPANIA DEL COMPANIA DE LA COMPANIA DEL COMPANIA DE LA COMPANIA DEL COMPANIA DE LA COMPANIA DEL COMPANIA DE LA COMPANIA DEL COMPANIA DEL COMPANIA DE LA COMPANIA DE LA COMPANIA DEL COMP
	Ala CaraVal-Asp-Pro),
16	4(Asn-Ile-Thr-Val-Thr-Ala-Ser-Val-Asp-Pro),
17	8(Thr-Ala-Ser-Val-Asp-Pro-Val-IIe-Asp-Leu),
	12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
18	15 (19-A sp-Leu-Leu-Gin-Ala-Asp-Gly-Asn-Ala),
19	20/Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
20	26(Pro-Ser-Ala-Val-Lys-Leu-Ala-Tyr-Ser-Pro),
21	20(PTU-50: 1

	72(Leu-Asn-Ser-Thr-Val-Gin-Met-Pro-Ile-Ser),
1	72(Leu-Asn-Sci-III)
2	78(Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Glu-Phe),
	87(Gin-Vai-Leu-Ser-Thr-Thr-Ala-Lys-Giu-Phe),
3	Ala-Dm-Thr-Ala-Gly-Ash-1907
4	- Cly-Val-Val-Ser-Leu-Val,
5	133(Gly-Asn-Tyr-Ser-City VIII) peptides containing CFA/I pilus protein B-cell (antibody) epitopes (Starting
6	
7	Sequence # given)
	- il enitones
,	CFA/I pilus protein B-cell epitopes
8	3(Lys-Ana-Ile-Thr-Val-Thr-Ala-Ser-Val),
9	3(Lys-Ana-ne and a specific and a sp
10	Alas Jeu-Pro-Set-Ala-Val),
11	22(Gly-Asn-Ala-Lee - 1) 32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-Thr-Phe-
12	
13	Glu-Ser-Tyr-Arg-Val),
-	32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe),
14	29.7 vs-Thr-Phe-Glu-Ser-Tyr-Arg-Val),
15	66(Pro-Gln-Leu-Thr-Asp-Val-Leu-Asn-Ser),
. 16	93(Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala),
17	93(Ala-Lys-Clu-7 no Clu Thr-Ala-Pro-Thr),
18	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr),
	124(Lys-Int-Ala-Ory-Ala-Gly-Asn-Tyr-Ser), and 127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser)
19	127(Gly-Thr-Ala-Fio Thr-Ala-Pro-Thr-Ala-Gly-Asn- Tyr- 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn- Tyr-
20	Ser), and mixtures thereof.
21	Scr), and management of the script of the sc

	(12) synthetic peptides containing CFA/I pilus protein T-cell and
1	(12) synthetic pepader
2	B-cell (antibody) epitopes (Starting Sequence # given)
2	CFA/I pilus protein T-cell epitopes
3 .	3(Lys-Ana-Ile-Thr-Val-Thr-Ala-Ser-Val),
4	11(Val-Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
5	22(Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
6	22(Gly-Asn-Ala-Zer-Lys-Thr-Phe-Lys-Thr-Phe- 32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-Thr-Phe-
7	
8	Glu-Ser-Tyr-Arg-Val),
	32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe),
9	38(Lys-Thr-Phe-Glu-Ser-Tyr-Arg-Val),
10	66(Pro-Gln-Leu-Thr-Asp-Val-Leu-Asn-Ser),
11	93(Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala),
12	93(Ala-Lys-Glu-File Growth Ala-Pro-Thr)
13	124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr),
	127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
14	127(Gly-Thr. 700) 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-
15	
16	Ser); and synthetic peptides containing CFA/I pilus protein T-cell and B-cell (antibody)
17	synthetic peptides containing Caracapa
18	epitopes (Starting Sequence # given)
	CFA/I pilus protein B-cell epitopes
19	3(Lys-Asn-Ile-Thr-Val-Thr-Ala-Ser-Bal-Asp-Pro),
20	3(Lys-Asir-lic 112) 8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-IIe-Asp-Leu-Leu-Gln-
21	
22	Ala-Asp),

	11(Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
1	20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
2 .	20(Ala-Asp-Gly-Ash-Ala-Bro-Thr-Ala-Gly-Ash-Tyr-Ser), and 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Ash-Tyr-Ser)
3	124(Lys-Thr-Ala-Gly-Inr-Ala-Flo
4	126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and mixtures
5	thereof.
	thereof. We contemplate that the peptides can be used in vaccine constructed
6	for systemic administration.
7	for systemic administration of the systemic administration of
8	The peptides in (8), (9), and (10) above can be made by classical
9	The peptides in (b), (9), and (co)
10	solution phase synthesis, solid phase synthesis or recombinant DNA
11	solution phase synthesis, seem that technology. These peptides can be incorporated in an oral vaccine to prevent
12	infection by CFA/I bearing enteropathogenic E. coli.
	harrin offered examples provide methods for musually, wanted
13	any implied limitation, the practice of this invention in the prevention of
14	diseases caused by enteropathogenic organisms.
15	The profile of the representative experiments have been chosen to
16	The profile of the representational polymeric matrix-antigen
17	illustrate the effectiveness of the immunogenic polymeric matrix-antigen
18	composites.
	composites. All temperatures not otherwise indicated are in degrees Celcius (C)
19	and parts or percentages are given by weight.
20	MATERIALS AND METHODS
21	Animals. New Zealand White male rabbits were purchased from
22	Animals. New Zealante Hazelton Research Products (Denver, PA), and were shown to be free of
23	Hazelton Research Products (Deliver, 1997) swabs. Animals were 1-2 kg of
24	Hazelton Research Florida Animals were 1-2 kg of current RDEC-1 infection by culture of rectal swabs. Animals were 1-2 kg of

body weight and lacked agglutinating anti-AF/R1 serum antibody at the time of 1 Antigens. AF/R1 pili from E. coli RDEC-1 (015:H:K non-typzble) the study. 2 were purified by an ammonium sulfate precipitation method. The final 3 preparation migrated as a single band on SDS-polyacrylamide gel electrophoresis and was shown to be greater than 95% pure by scanning with 5 laser densitometry when stained with coomassie blue. Briefly, equal molar 6 parts of DL-lactide and glycolide were polymerized and then dissolved to 7 incorporate AF/R1 into spherical particles. The microspheres contained 0.62% 8 protein by weight and ranged in size from 1 to 12 micrometers. Both the 9 microencapsulated and non-encapsulated AF/R1 were sterilized by gamma 10 11 irradiation (0.3 megarads) before use. Synthetic peptides (16 amino acids each) were selected by theoretical 12 criteria from the amino acid sequence of AF/R1 as deduced from the 13 nucleotide sequence. Three sets of software were used for the sections. 14 Software designed to predict B cell epitopes based on hydrophilicity, 15 flexibility, and other criteria was developed by the University of Visconsin 16 Genetics Computer Group. Software designed to predict T cell eximpes was 17 based on the Rothbard method was written by Stephen Van Albes (The Walter 18 Reed Army Institute of Research, Washington, D.C.). Software sixigned to 19 predict T cell epitopes based on the Berzofsky method is publishes as the 20 AMPHI program. The selected peptides were synthesized by using 21 conventional Merrifield solid phase technology. AF/R1 40-55 22 (Thr-Asn-Ala-Cly-Thr-Asp-Ile-Gly-Ala-Asn-Lys-Ser-Phe-Thr-LenLys) was 23 24

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various dilutions of antigen and were incubated at 37°C in 5% CO₂. In other experiments, cultures were conducted in a 24-well plates. In these experiments, 5 x 10° cells were cultured with or without antigen in a 2 ml volume. After 4 days, 100 microliters aliquots of cells were transferred to 96-well plates for pulsing and harvesting. Previous experiments have demonstrated that optimal concentrations of antigen range from 150 ng/ml to 15 micrograms/ml in the 96-well plate assay and 1.5 ng/ml to 150 ng/ml in the 24-well plate assay. These were the concentrations employed in the current study. All cultures were pulsed with 1 Ci [3H]thymidine (25 Ci/mmol, Amersham, Arlington Heights, IL) on day 4 of culture and were harvested for scintillation counting 6 hours later. Statistics. All cultures were conducted in replicates of four, and standard deviations of the counts per minute (cpm) generally range from 5-15% of the average cpm. In experiments where comparison of individualanimals and groups of animals is desirable, data is shown as a stimulation index (SI) to facilitate the comparison. SI were calculated by dividing the mean of cultures with antigen by the mean of cultures without antigen (media 16 control). Statistical significance (p value) was determined by comparing the 17 maximum response for each antigen to the media control using the Statem's E 18 19 test. RESULTS 20 Lymphocyte proliferation in response to protein and peptife anigens 21 of AF/R1. To determine if lymphoid tissues from AF/R1 immunerations 22 respond in vitro to the antigens of AF/R1, the immunity in a rabbit with 23

preexisting high levels of anti-AF/R1 serum IgG was boosted twice by injection of 50 micrograms of purified AF/R1 pili i.p. seven days apart. A week after the final boost, in vitro lymphocyte proliferation of spleen and MLN cells demonstrated a remarkable response to AF/R1 pili. In response to the synthetic peptides, there was a small, but significant proliferation of the spleen cells to all the AF/R1 peptides tested as compared to cell cultures without antigen. Cells from the spleen and Peyer's patches of non-immune animals failed to respond to either AF/R1 or the synthetic peptides.

Microencapsulation of AF/R1 potentiates the mucosal cellular immune response. To evaluate the effect that microencapsulation of AF/R1 may have on the cellular mucosal immune response to that antigen, naive rabbits were primed twice with 50 micrograms of either microencapsulated or non-encapsulated AF/R1 by endoscopic intraduodenal inoculation seven days apart. All rabbits were monitored daily and showed no evidence of clinical illness or colonization by RDEC-1. One week following the last priming, the rabbits were sacrificed and lymphoid tissues were cultured in the presence of AF/R1 pili or peptide antigens. In rabbits which had received non-encapsulated AF/R1, Peyer's Patch cells demonstrated a low level but significant proliferation in vitro in response to AF/R1 pili (Fig.13), but not to any of the AF/R1 synthetic peptides (Fig.14a-d). However, in rabbits which had received microencapsulated AF/R1, Peyer's Patch cells demonstrated a markedly enhanced response not only to AF/R1 (Fig.13 but now aspossed to the AF/R1 synthetic peptides 40-55 and 79-94 (Fig.14 a and 45). Institute.

one of two rabbits primed with microencapsulated AF/R1 (rabbit 135) 1 responded to AF/R1 108-123, but not AF/R1 40-47/79-86 (Fig14c and 4d). 2 In contrast, the other rabbit in the group (rabbit 134) responded to AF/R1 3 40-47/79-86, but not to AF/R1 108-123 (Figl 4 d and Lac). Response of MLN cells to the antigens of AF/R1. Studies have shown 5 that cells undergoing blastogenesis in the MLN also tend to home into mucosal 6 areas, but experiments requiring in vitro lymphocyte proliferation of rabbit 7 MLN cells are difficult to conduct and to interpret due to non-specific high 8 background cpm in the media controls. Our studies have shown that this 9 problem can be avoided by conducting the proliferative studies in 24-well 10 plates, and then moving aliquots of cells into 96-well plates for pulsing with [3H]thymidine as described in materials and methods. This method of culture 11 was employed for the remainder of the studies. The MLN cells of all rabbits 12 13 demonstrated a significant proliferation in vitro in response to AF/R1 pili 14 regardless of whether they had been immunized with microencapsulated or 15 However, only the rabbits which had non-encapsulated AF/R1. 16 received microencapsulated AF/R1 were able to respond to the AF/R1 17 synthetic peptide 40-55 (Fig.19'). The MLN cells of rabbit 134 also responded to AF/R1 79-94 (p<0.0001), AF/R1 108-123 (p<0.0001), and 18 19 AF/R1 40-47/79-86 (p=0.0004); however, none of the other rabbits 20 demonstrated a MLN response to those three peptides (data not shown). 21 Response of spleen cells to the antigens of AF/R1. Proliferative responses of spleen cells to AF/R1 were very weak in all animals tested (data 22 not shown). However, in results which paralleled the responses in MIN cells, 23

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there was a significant response to AF/R1 40-55 in rabbits which had been primed with microencapsulated AF/R1 (Fig. 20'). There was no response to the other AF/R1 synthetic peptides by spleen cells in either group of animals. The weak response of spleen cells to AF/R1 provides further evidence that these animals were naive to AF/R1 before the study began, and indicates that the observed responses were not due to non-specific stimulative factors such as lipopolysaccharide.

SUMMARY

We have shown that there is an enhanced in vitro proliferative response to both protein and its peptide antigens by rabbit Peyer's patch cells following intraduodenal inoculation of antigen which had been homogeneously dispersed into the polymeric matrix of biodegradable, biocompatible microspheres. The immunopotentiating effect of encapsulating purified AF/RI pili as a mucosal delivery system may be explained by one or more of the following mechanisms: (a) Microencapsulation may help to protect the antigen from degradation by digestive enzymes in the intestinal lumen. (b) Microencapsulation has been found to effectively enhance the delivery of a high concentration of antigen specifically into the Peyer's patch. (c) Once inside the Peyer's patch, microencapsulation appears to facilitate the rapid phagocytosis of the antigen by macrophages, and the microspheres which are 5-10 micrometers become localized within the Peyer's patch. (d) Microencapsulation of the antigen may improve the efficiency of antigen presentations by decreasing the amount of enzymatic degradation that takes place inside the macrophage before the epitopes are protected by combining with

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Class II major histocompatibility complex (MHC) molecules. (e) The slow,
factions may produce a depot effect and the
retention of antigen by the follicular dendritic cell. (f) If the antigen of interest
is soluble, microencapsulation changes the antigen into a particulate form
is soluble, microencapsulation changes are all response by shifting the
which appears to assist in producing an IgA B cell response by shifting the
which appears to assist in process which appears to assist in process cellular immune response towards the T _H and thereby not encouraging a cellular immune response towards the T _H and thereby not encouraging a
There is evidence that the GALI may
microbial and non-microbial (1008) 2.12
site entiren when it is first encountered, and that
the special antigenic characteristics that make
s adjactizens, but they are antigenic because
the particulate nature of the particulate nature
may serve to mimic that context. It may be important to note that we also
observed a significant response to AF/R1 in animals inoculated with
observed a significant response to 712 of this antigen which was still in its native non-encapsulated pili; thus, some of this antigen which was still in its native
non-encapsulated pili; thus, some of unit artigory
non-encapsulated pill; titus, some some some some some some some some
form was able to enter the 1990 1990 that AF/R1 is known to mediate the attachment of RDEC-1 to the Peyer's
as the antigen employed in this type of study will
Macella, one would expect to see an even grant
attach to micrometer in the responses of animals which had received microencapsulated
and antigen.
there used in these experiments included a
The 1 to 5 micrometer particles have been aloue
from 1 to 12 micrometers. The state of the 12 micrometers, the to disseminate to the MLN and spleen within migrating macrophages, thus, the
to disseminate to the MLN and specific

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observed proliferative responses by cells from the MLN and spleen may reflect priming of MLN or splenic lymphocytes by antigen-presenting/accessory cells which have phagocytosed 1 to 5 micrometer antigen-laden microspheres in the Peyer's patch and then disseminated onto the MLN. Alternatively, these responses may be a result of the normal migration of antigen stimulated lymphocytes that occurs from the Peyer's patch to the MLN and on into the general circulation before homing to mucosal sites. Proliferative responses by MLN cells are of interest because it has been shown that cells undergoing blastogenesis in the MLN tend to migrate onto mucosal areas. However, studies involving in vitro lymphocyte proliferation of rabbit MLN cells can be very difficult to conduct and to interpret due to non-specific high, background cpm in the media controls. By simultaneously conducting experiments using different protocols, we have found that this problem can be prevented by avoiding the use of fetal calf serum in the culture and by initially plating the cells in 24-well plates. Using this method, the blasting lymphocytes are easily transferred to a 96-well plate where they receive the [3H]thymidine, while fibroblasts and other adherent cells remain behind and thus do not inflate the background cpm.

The proliferative response to the peptide antigens was of particular interest in these studies. The rabbits that received non-encapsulated AF/R1 failed to respond to any of the peptides tested either at the level of the Peyer's patch, the MLN, or the spleen. In contrast, Peyer's patch cells from the animals that received microencapsulated AF/R1 responded to all the peptides tested with two exceptions: Rabbit 134 did not respond to AF/R1 108-123,

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and rabbit 135 did not respond to AF/R1 40-47/79-86. The reason for these non-responses is not clear, but it probably is not due to MHC restrictions as evidenced by the fact that rabbit 134 was able to respond to AF/RI 108-123 at the level of the MLN. The non-responses may be due to varing kinetics of sensitized T cell migration in different rabbits, or they may reflect differences in the efficiency of antigen presentation by cells from different lymphoid tissues of these animals. Of all the synthetic peptides tested, only AF/R1 40-55, (the one selected as a probable B cell epitope), was recognized by serum from an AF/R1 hyperimmune rabbit. In addition, this peptide was the only one that was uniformly recognized by Peyer's patch, MLN, and spleen cells from both rabbit. In addition, this peptide was the only one that was uniformly recognized by Peyer's patch, MLN, and spleen cells from both rabbits that were immunized with microencapsulated AF/R1. The recognition by anti-AF/R1 serum antibodies indicates that the amino acid sequence of this peptide includes an immunodominant B cell epitope. Thus AF/R1 40-55 may readily bind to antigen-specific B cells thereby leading to an efficient B cell presentation of this antigen to sensitized T cells. Even though AF/R1 40-55 was not selected as a probable T cell epitope by either the Rothbard or Berzofsky methods, the current study clearly indicates that this peptide can also stimulate a proliferative immune response. Although further studies are required to definitively show that the proliferating cells are indeed T cells, the responses observed in this study are most likely due to the blast transformation of cells from the lineage. Therefore, AF/R1 40-55 appears to contain a T cell epitope in addition to the immunodominant B cell epitope, and this area of the

	important role in the overall immune
1	AF/R1 protein may thereby play an important role in the overall immune
2	response and subsequent protection against RDEC-1.
3	or serious segronses of spleen cells was lot a
	and the this may be simply a matter of the answer
4	mba mbbits in this study were satisfied
5	to antigen. This relatively short since
6	after their first exposure was aligned. have provided sufficient time for cells that were produced by Peyer's patch and have provided sufficient time for cells that were produced by Peyer's patch and
7	have provided sufficient time for centre and sufficient numbers. MLN blasts to have migrated as far as the spleen in sufficient numbers.
8	MLN blasts to have migrated as far as the specific would not only assist in the
9	MLN blasts to have inigrated. An ideal mucosal vaccine preparation would not only assist in the
10	uptake and presentation of the immunogen of interst, but it would also be
11	carrier molecules or adjuvants with
	in a production of delay regulatory approvat.
12	innerpheres appears to provide an ideal indecode
13	immunogens because the observed immunogens
14	system for oral vaccine initiatings effect is achieved without the need for carriers of adjuvants. This ability may
15	prove to be of great value, particularly to enhance the delivery of ord
16	prove to be of great value, particularly to
17	synthetic peptide vaccines to the GALT.
	and the Monkeys
10	TABLE 10 Linear B-Cell Epitopes of CFA/I in Monkeys
18	
	Sequence Individuals
19	Consensus Site
20	Position Responding Communication
	VDPVIDLLQ
21	1. 11-21 3

PCT/US98/01556 WO 98/32427

1 2 3 4 5 6	 93-101 124-136 66-74 22-29 32-40 38-45 3-11 	2 2 2 2 1 1	GPAPT PQLTDVLN GNALPSAV KTF*
8 9 ,	•Overlap betw	reen epit	ope 6 and 7

^{*}Overlap between epitope 6 and 7

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		TABLE 11	
1	Prediction of T.S	ell epitopes within	the CFA/I molecule
2	Predicted Ampl	nipathic Segments	Rothbard Criteria
3	7 az blocks	11 22 blocks	16
5	22-25	8-11	
	34-38	32-44	30
6	40-46	51-71	38
7	50-53	86-92	44
8	56-62	102-108	57
9 '	64-71	130-131	61
10		135-137	70
11	104-108		116
12	131-137		124
13			127
14			137
15			

The sequence numbers of the first amino acid of the predicted T cell epitopes are shown. Software designed to predict T cell epitopes based on the Berzofsky method was published as the AMPHI program. It predicts amphipathic amino acid segments by evaluating 7 or 11 residues as a block and assigning a score to the middle residue of that block. Software designed to predict T cell epitopes based on the Rothbard method was written by Stephen Van Albert (The Walter Reed Army Institute of Research, Washington, D.C.).

22 Van Albeit (Tille TABLE 11 23

•	156
, 1	Amino acid sequence of immunodominant T cell epitopes*
2	Residue
3	Numbers Amino Acids
	8 17 Thr Ala Ser Val Asp Pro Val Ile Asp Leu
4	40-49 Phe Glu Ser Tyr Arg Val Met Thr Gln Val
5	72 81 Leu Asn Ser Thr Val Gln Met Pro Ile Ser
6	Ser Gly Val Val Ser Leu Val Met
7	that supported a significant pro-
8 ,	serine at either position 2, 3, of 4, mind a
9	Some of the most robust response
10	to entire residue at the unit possession
11	peptides that contained a serific term peptides that contained term peptides that contai
12	1.00
13	T cell epitopes is shown. DEMONSTRATIVE EVIDENCE OF PROTECTIVE IMMUNITY DEMONSTRATIVE EVIDENCE OF PROTECTIVE IMMUNITY
14	etaroadherent diarrhea producing E. Con
15	is by the adhesin (AF/R1 pill). The adhesia
16	It may intitiate a mucosal response
. 17	the gut The incorporation of
18	respectible microspheres enhances most
19	respones to RDEC-1. We have demonstrated that immunization with AF/RI
20	respones to RDEC-1. We have been respones to RDEC-1. Pili in microspheres protect rabbits against infection with RDEC-1. Pili in microspheres protect rabbits against infection with RDEC-1.
21	Pili in microspheres protect labora ega- Pili in microspheres protect labora ega- Six rabbits received intra-duodenal immunization of AF/R1
22	Six rabbits received intra-discostant at 200 ug AF/R1 on day then microspheres (0.62% coreloading by weight) at 200 ug AF/R1 on day then
23	microspheres (0.62% coreloading by weight) and 21 fallowed boosted with 100 ug AF/R1 in microspheres on days 7, 14, and 21 fallowed
24	boosted with 100 ug AF/R1 in microsyncies

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by RDEC-1 challenge with 10⁴ organisms one week latter than observed for 1 week and then sacrificed, unimmunized rabbits were challenged with 10^s RDEC-1 only and observed 1 week than sacrified. Also, 2 rabbits were immunized only then were sacrificed 10 days latter. Only one of these animals had bile IgA antibodies to AF/R1. but both had specific sensitized T cells which released IL-4 upon challenge in the spleen, Peyer's patch and illeal lamina propria. All nine immunized animals developed diarrhea and weight loss which was significant at the p < .001 level compared to the immunized animals which displayed no diarrhea and no weight loss. The immunized animals colonized the intestinal tract with RDEC-1 the same as the unimmunized animals. However, there was a striking difference regarding the adherence of RDEC-1 to the mucosa. No adherence was seen in cecum in the immunized animals compared to 4/7 in the unimmunized side animals. This difference was significant to the p < .01 level. The RDEC-1 exposure although not producing disease in the immunized animals did effect a booster immunization as relected in the increase in anti-AF/R1 antibody containing cells in the muscosa similiar to the immunized rabbits. This study clearly demonstrated complete protection against RDEC-1 infection and strongly indicates similiar results should be expected with entertoxigenicity E. of using the Colony Forming Antigens (CFA's) in microsphere vaccines. SUMMARY STATEMENT OF PROTECTIVE IMMUNITY SHOWINGS RDEC-1 infection of rabbits causes an enteroadherent E. coli marrheal disease, and provides a model for the study of adherence-factor immunity. Pilus adhesions are vaccine candidates, but purified pili are subject to intestinal

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degradation. Previously we showed potentiation of the mucosal cellular · immune response to the AF/R1 pilus of RDEC-1 by incorporation into biodegradable polylactide-coglycolide microspheres (AF/R1-MS). We now present efficacy testing of this vaccine. Six rabbits were primed with 200 ug and boosted with 100 ug of AF/R1-MS weekly x3, then challenged at week 5 with 10° CFU of RDEC-1 expressing AF/R1. Nine unvaccinated rabbits were also challenged. Two rabbits vaccinated with AF/R1-MS were sacrificed at week 5, without challenge, for measurement of anti-AF/R1 antibodies in bile (by ELISA) and anti-AF/R1 containing cells (ACC) in the intestinal lamina propria (by immunohistochemistry). Attachment of RDEC-1 to intestinal epithelial cells was estimated (0.4+) by immunoperoxidase staining of histologic sections. Colonization of intestinal fluid was measured by culture of intestinal flushes. Results: Rabbits given AF/R1-MS remained well and 4/6 gained weight after challenge, whereas 9/9 unvaccinated rabbits lost weight after challenge (mean weight change +10 vs -270 gms p < .001), (see Figure 35). The mean score of RDEC-1 attachment to the cecal epithelium was 0 in vaccinated, and 2+ in unvaccinated animals (see Figure 36). RDEC-1 colonization (log CFU/gm) in cecal fluids was similar in both groups (mean 6.3 vs 7.3; p=.09) (see Figure 34). ACC were not seen in the lamma propria of vaccinated but unchallenged animals, but anti-pilus IgA antibody levels in bile were increased 1 S.D. over negative controls in 1 animal. Condusions: Vaccination with AF/R1-MS was safe and protected rabbits against EDEC-1 disease. Protection was associated with interference with RDEC-1 atherence to the mucosal surface, but lumenal colonization was not prevented.

More recently, applicants have focused on areas of this invention related to an immunostimulating composition for the burst-free, sustained, programmable release of active material(s) 1 4 over a period from 1 to 100 days, which comprises encapsulating 5 microspheres, which may contain a pharmaceutically-acceptable 6 adjuvant, wherein said microspheres are comprised of (a) a blend 7 of uncapped and end-capped biodegradable-biocompatible poly(DLlactide-co-glycolide) as the bulk matrix, wherein the relative 9 ratio between the amount of lactide and glycolide components are 10 within the range of 90:10 to 40:60 and the poly(DL-lactide-co-11 glycolide) is a blend of uncapped and end-capped forms in ratios 12 ranging from 100:0 to 1 to 99, and (b) active material such as an 13 immunogenic substance comprising Colony Factor Antigen (DFA/II, 14 hepatitis B surface antigen (HBsAg)), and/or a physiologically 15 similar antigen that serves to elicit the production of 16 antibodies in a mammal (human or nonhuman). These areas of invention are referred to hereim after 18 as Part II and Part III, respectively, and are itemized as An immunostimulating composition for the burst-free, 19 follows: 21 sustained, programmable release of active material(s) over a 22 period from 1 to 100 days, which comprises encapsulating-23 microspheres, which may contain a pharmaceutically-acceptable 24 adjuvant, wherein said microspheres having a diameter between 1 25 nanogram (ng) to 10 microns (um) are comprised of (a) a blumd of 26 uncapped and end-capped biodegradable-biocompatible poly [15-27 lactide-co-glycolide) as the bulk matrix, wherein the relative 28 ratio between the amount of lactide and glycolide components are 29 within the range of 90:10 to 40:60, and the poly(DL-lactide-co-30 glycolide) is a blend of uncapped and end-capped forms in ratios 31 ranging from 100:0 to 1 to 99, and (b) active material such as an 32 immunogenic substance comprising Colony Factor Antigen (CTA/II), 33 hepatitis B surface antigen (HBsAg), and/or a physiologically 34 similar antigen that serves to elicit the production of 35 antibodies in a mammal (human or nonhuman).

1 2 3	 An immunostimulating composition according to Item 1 wherein the amount of said immunogenic substance is within the range of 0.1 to 1.5% based on the volume of said bulk matrix. An immunostimulating composition according to Item 2 wherein the relative ratio between the lactide and glycolide component is within the range of 48:52 to .52:48
7 8 9	4. An immunostimulating composition according to Item. 2 wherein the size of more than 50% of said microspheres is between 5 to 10 um in trameter by volume.
10 11	5. A vaccine comprising an immunostimulating composition of Item 4 and a sterile, pharmaceutically-acceptable carrier therefor.
12	6. A vaccine comprising an immunostimulating composition of Ttem-5 wherein said immunogenic substance is Colony Factor Antigen (CFA/II).
14 15	7. A vaccine comprising an immunostimulating composition of Ite 5 wherein said immunogenic substance is hepatitis B surface antigen (HBsAg).
16 17 18	8. A method for the vaccination against bacterial infection comprising administering to a human, an antibacteric idally effective amount of a composition of Item 6.

1	9. A method according to I tem 7 wherein the bacterial infection is caused by
2	a bacteria selected from the group consisting essentially of Salmonella typhi,
3	Shigella Sonnei, Shigella Flexneri, Shigella dysenteriae, Shigella boydii,
4	Escheria coli, Yibrio cholera, yersinia, staphylococus, clostridium, and
-	campylobacter.
5	CAMPATOURE .
6	10. A method for the vaccination against viral infection comprising
7	administering to a human an antivirally effective amount of a composition of
8	'Item 7.
9	11. A diagnostic assay for bacterial infections comprising a composition of
10	Item 4.
•	infections
11	12. A method of preparing an immunotherapeutic agent against infections
12	caused by a bacteria comprising the step of immunizing a plasma donor with a
13	vaccine according to . Item 6 such that a hyperimmune globulin is produced
14	which contains antibodies directed against the bacteria.
	•
15	13. A method preparing an immunotherapeutic agent against infections caused
	by a virus comprising the step of immunizing a plasma donor with a vaccine
16	according to Item. 7 such that hyperimmune globulin is produced which
17	contains antibodies directed against the hepatitis B virus.
18	COUNTING STIFFOOTION CONTINUES CONTI

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1	14. An immunotherapy method comprising the step of administering to a
2	14. An immunomerapy means subject an immunostimulatory amount of hyperimmune globulin prepared
3	according to Item 12.
4 5 6	15. An immunotherapy method comprising the step of administering to a subject an immunostimulatory amount of hyperimmune globulin prepared according to Lem 13.
7 8	16. A method for the protection against infection of a mammal (human or nonhuman animal) by enteropathogenic organisms or hepatitis B virus
9	comprising administering to said mammal an immunogenic amount of the
10	immunostimulating composition of Item 3.
11	17. A method according to Item 16 wherein the immunostimulating composition is administered orally.
12	
13	18. A method according to Item 16 wherein the immunostimulating
14	composition is administered parenterally.
15	PART II In sum, the Colony Factor Antigen (CFA/II) from enterotoxigmic E
16	In sum, the Colony Factor Paraget (Soli (ETEC) prepared under GMP was successfully incorporated into
17	coli (ETEC) prepared under GMP was successfully under GMP and found to biodegradable polymer microspheres (CFA/II BPM) under GMP and found to
18	biodegradable polymer microspheres (crass and intra-duodenally to rabbits.
19	biodegradable polymer new poly
. 20	CFA/II was incorporated into poly (D, L-122222 2 2 2

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microspheres which were administered by direct endoscopy into the duodenum. Following vaccination, Peyer's patchcells responded by lymphocyte proliferation to in vitro challenge with CFA/II indicating the CFA/II BPM to be immunogenic when administered intra-intestinally. Also, B cells secreting specific anti CFA/II antibodies were found in spleens following vaccination. No pathological changes were found following total necropsies of 10 rabbits vaccinated with CFA/II BPM. As a potency test, high serum IgG antibody titers to CFA/II were produced following intra- muscular administration of CFA/II BPM to additional rabbits. The CFA/II BPM contained 63% between 5-10 um by volume particle size distribution; 1.17% protein content; 2.15% moisture; < .01% acetonitrile; 1.6% heptane; 22 nonpathogenic bacteria and 3 fungi per 1 mgm protein dose; and passed the general safety test. We conclude that the CFA/II BPM oral vaccine is immunogenic and safe to begin a Phase I clinical safety study following IND approval. INTRODUCTION Enterotoxigenic Escherichia coli (ETEC) causes diarrheal disease with an estimated 650,000,000 cases anually in developing countries resulting in 500,000 deaths predominantly in the pediatric age groups. Currently there is no vaccine against ETEC induced diarrhea. The availability of an effective oral vaccine would be of great value to the people of South America, Africa and and Asia as well as the millions of people who travel to these high risk areas and account for half of the annual cases. 21

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	The first step in pathogenesis is adherence to the small intestine
L	The first step in pathogenesis is a called colonization factor epithelial cells by protein fimbrial (pilus) adhesins called colonization factor epithelial cells by protein fimbrial (pilus) adhesins called colonization factor
2	epithelial cells by protein fimbriai (plus) been recognized, CFA/I, CFA/II and antigen (CFA). Three major CFAs have been recognized, CFA/I, CFA/II and
3	antigen (CFA). Three major CFAs have so
4	CFA/IV. (25)

Ten human volunteers who were immunized orally twice weekly for 4 weeks with CFA/II developed a poor antibody response and did not show any significant protection when challenged with pathogenic ETEC (26). This disappointing response was attributed to adverse effects of gastric acid, even at neutral pH, of fimbrial proteins (27). When the vaccine was administered by inoculation directly into the duodenum, 4 of 5 immunized volunteers developed a significant rise in secretory IgA with CFA/II antibody (26).

D and L-lactic acid and glycolic acid, as homo- and copolymers, are biodegradable and permit slow and continued release of antigen with a resultant adjuvant activity. These polymers have · been shown to be safe in a variety of applications in human beings and in animals (28-32). Delivery of antigens via microspheres composed of biodegradable, biocompatible lactide/glycolide polymers (29-32) may enhance the mucosal response be protecting the antigen from digestion and targeting them to lymphoid cells in Peyer's patches (29-32). McQueen et al. (33) have shown that E con AF/R! pili in PLGA microspheres, introduced intra-duodenally in rabbits, protected them against diarrhea and weight loss when challenged with the parent strain

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rabbit diarrheagenic strain of E coli (RDEC-1). Only one vaccinated rabbit of
rabbit diarrheagenic strain of a secontrast, all control
to the one had soft perfected see
rabbit diarrheagenic strain of E control six lost weight and only one had soft pelleted stool. In contrast, all control
herame ill, lost weight, —
six lost weight and only one included weight, and shed soft pellets unvaccinated animals became ill, lost weight, and shed soft pellets or unformed mucoid stool. Significant lymphocyte proliferation to
Significant lymphosy
or unformed masses
or unformed mucoid stool. Significantly IgA anti AF/R1 antibody AF/R1 from Peyer's patches and ordinary IgA anti AF/R1 antibody
AFIRE
levels were seen.
10.4

In order to improve the CFA/II vaccine it was incorporated into PLGA microspheres under GMP in order to protect it from digestion and target it to the intestinal lymphoid system. The CFA/II BPM vaccine has undergone pre-clinical evaluation and has been found to be safe and immunogenic.

MATERIALS AND METHODS

Preparation of CFA/II Pilus Vaccine. Under Good Laboratory and Good Manufacturing Practices, E. coli. strain M424C1-06;816 producing CFA/II were cultured in 75-80 CFA agar plates (24 x 24 cm) for 24 hrs then harvested by scraping. The harvest was homogenized at slow speed for 30 minutes with over head drive unit and cup immersed in an ice bath. The homogenate was centrifuge at 4° C at 16, 500 x g for 30 minutes. The supernatant saved and the pellet rehomogenized and centrifuged with the supernatants pooled. The supernatant pool was centrifuged at 50,000 x g for 45 minutes. The supernatant treated with ammonium sulfate at 20% satuaration, stirred 30 minutes at 4° C than stored at 4° C for 16 hrs then

100 sayed and treated
centrifuged at 19,700 x g for 30 minutes. The supernatant saved and treated
centrifuged at 19,700 x g for 30 minutes. with ammonium sulfate at 45% saturation, stirred 30 minutes at 4° C, stored at with ammonium sulfate at 45% saturation, stirred 30 minutes. The pellet
The same of the sa
with ammonium sulfate at 43 70 salurants. The peller
the centifuged at 27
4°C for 60°C Tornauli and likely
4. C for 66-72 hrs, then centrifuged at 257 was resuspended in about 100 mls of PBS containing 0.5% formalin and held was resuspended in about 100 mls of PBS containing 0.5% formalin and held
was resuspended at a contract PBS at 4. C using a total
was resuspended in about 100 mis of 120 or against PBS at 4. C using a total at 22 for 18 hrs then dialyzed for 45-50 hrs against PBS at 4. C using a total
The dialysis was
of 12 liters in 2 liter and fuchsin
- colio lising incide
of 12 liters in 2 liter amounts. The contained less then 0.03% formalin using Nessler's reagent and fuchsin contained less then 0.03% formalin using Nessler's reagent and fuchsin sulfuose acid reagent. The final product contained 1 mgm protein/ml PBS,
The final product contained I mg. P
sulfuose acid reagent. The inner pro-
safety test.
was sterile and passed the general safety test. Was sterile and passed the general safety test. Two ml of the CFA/II
was store. Two mi of the Critical

Preparation of Desalted CFA/II Vaccine. Two ml of the CFA/II vaccine were placed into a Centricon 30 tube and centrifuged at 1700 rpm at 4-6. C (Beckman model GPR centrifuge equipped with GA-24fixed angle rotor) until all the buffer solution passed through the filter (about 90-120 minutes). Sterile water was added to each tube to disperse the CFA/II retained on the filter. The desalted antigen dispersions from all tube were pooled and then divided into five equal parts by weight so as to contain 20 mg of the CFA/II divided into five equal parts by weight so as to contain 20 mg of the CFA/II each. The desalted antigen dispersion was stored at -10 to -20° C.

Freeze Drying of the Desalted CFA/II Dispersion. 80 mg of sucrose was added to each part of the CFA/II dispersion. The resulting misture was flash-frozen using a dry ice-acetone bath (100-150 ml od acetone and 50-100 g of dry ice). The frozen solution was freeze dried overnight using Repp Sublimator 16 freeze dryer at vacuum of 1 micrometer of mercury and a shrift temperature not exceeding 37°C.

CFA/II Biodegradable Polymer Microspheres. Particle size distribution. About 1 mgm of microspheres were 1 dispersed in 2 ml of 1% Polysorbate 60 (Ruger Chemical Co. Inc. Irvington, 2 N.J.) in water in a 5 ml capacity glass vial by sonication. This dispersion was 3 observed under a calibrated optical microscope with 43x magification. Using a precalibrated eye-piece micrometer, the diameter of 150 randomly chosen 5 microspheres, was determined and the microsphere size distribution was 6 7 determined Scanning Electron Microscopic Analysis. Microspheres were 8 sprinkled or the surface of 10mm stub covered with a non-conductive adhesive 9 (Sticky-Tab, Ernest F. Fullem, Inc., Lutham, N.Y.) Samples were coated 10 with gold/palladium in an automatic sputter-coating opparatus (Samsputter-2A, 11 Tonsimis Research Corporation). The samples were examined with a Hiuchi 12 S-450 scanning electron microscope operated at 15-20 KV. 13 Preparation of CFA/II Microspheres. Solvent extraction techique was 14 used to encapsulate the freeze dried CFA/II into poly(lactide-co-15 glycolide)(Medisorb Techologies International, visocity 0.73 dl/g) 16 microspheres in the 1-10 um size-range to achieve theoretical antigen loading 17 of 1% by weight. The freeze dried antigen-sugar & matrix was dispursed in 18 an acetolnitrile solution of the polymer and then emulsified to achieve desired 19 droplet size. Microspheres were solidified and recovered by using beptane as 20 extracting solvent. The microsphere batches were pooled and vacuum dried to 21 22 remove traces of solvent.

	168 dissolved in 0.9%
	Protein Content. The CFA/II microspheres were dissolved in 0.9%
1	Protein Content. The CFA/II microsper SDS in O.1N NaOH for 18 hr with stirring then neutralized to pH 7 SDS in O.1N NaOH for 18 hr with stirring then neutralized with
2	SDS in O.1N NaOH for 18 hr with stirring their method was utilized with and assayed. The micro bicichoninic acid (BCA) method was utilized with
	and assayed. The micro bicicions
3	and assayed. The micro bicicional blanks and compared to both lactic acid and glycolic acid blanks and compared to
4	both lactic acid and glycolic acid blanks and compa- both lactic acid and glycolic acid blanks and results expressed as percent by bovine serum albumin (BSA) standard and results expressed as percent by
5	bovine 2.50 of CFA/II
6	weight. Moisture Content. One hundred and fifty mgm of CFA/II Moisture Content. One hundred and fifty mgm of CFA/II
7	Moisture Content. One hundred and firty may Moisture Content. One hundred and firty may microspheres were dissolved in 3 ml of acetonitrile by sonication for 3 hours. microspheres were dissolved in 3 ml of acetonitrile by sonication for 3 hours.
8	microspheres were dissolved in the microspheres wer
	observed was recorded and acetonitrile blank was substracted to determined
9	observed was recorded and acetonitine or
10	percent water content. Periduals, Ten mgm of CFA II
11	Acetonitrile and Heptane Residuals. Ten mgm of CFA II Acetonitrile and Heptane Residuals. Ten mgm of CFA II
12	Acetonitrile and Heptane Residuals Acetonitrile and Heptane Residuals microspheres were dissolved in 1 ml DMF then analysed using gas. microspheres were dissolved in 1 ml DMF then analysed using gas.
13	microspheres were dissolved in 1 ml DMP their description of blank microspheres.
14	chromatography and comparing peak heights to chromatography and
	acetonrile or heptane diluted in by weight.
15	The results are expressed as percent by weight.
16	The CEA/II microspherings
	Microbial load. One hundred mgm of CFA/II microsphere(single)
17	Microbial load. One hundred mgm of each of sterile saline than poured into 2 bind agas dose) are suspended in 2 ml of sterile saline than poured into 2 bind agas.
18	dose) are suspended in 2 ml of sterile saline under described after (\$ keeps in plates (1 ml each). All colonies are counted and identified after (\$ keeps in plates (1 ml each).
1	plates (1 ml each). All colonies are counted plates (1 ml each). All colonies are counted culture at 37° C and expressed as total number. Similiar amount of culture at 37° C and expressed as total number onto 4 different fungious
	culture at 37° C and expressed as and onto 4 different fungionalist
7	microspheres is in 0.25 mi angles.
	21 (Sabhiragar, casein peptone agar with Chiocartherical alars)
	plates (Sabhiragar, casein peptone agar with chloramphenol and genimycin or chloramphenol and genimycin or chloramphenol and genimycin or chloramphenol and genimycin or chloramphenol
	23 infusion agar with one

		table colories counted & identified and
		and cultured at 30° for 5 weeks and the colories counted & identified and
1		
_		expressed as total number. Canada Thirty mgm samples in
2		Minmenhere Silver
3		CFA/II Release From Microsopher Control of the Cont
J		placed in 2 ml conical upright microcarding
4		triplicate were placed in 2 ml content appropriate triplicate were placed in 2 ml content appropriate and kept immerized in a water 1 ml of PBS at pH 7.4. The tubes were capped and kept immerized in a water
		t of DRS at pH 7.4. The tubes were capped and
5		
		both maintained at 37° C with constant agree
6		intervals by control
7		at 1, 3, 6, 8, 15 and 22 nour times
,		and of Delicit top
8		at 1, 3, 6, 8, 15 and 22 hour time means at 1, 3, 6, 8, 15 and 22 hour time means at 1, 3, 6, 8, 15 and 22 hour time means at 1, 3, 6, 8, 15 and 22 hour time means that 1, 15 and 22 hour time means that 1, 15 and 22 hour time means that 1,
_		a sum nylor server
9		medium was collected through a 5 cm. medium was
		using the micro BCA method and companie
10		cumulative release of
		expressing results as percent cumulative release of CFA/II. General Safety Test. Two doses of one hundred mgm CFA/II General Safety Test. Two doses of one hundred mgm CFA/II
11		- Two does of one
12		General Salety 1991
		General Safety Test. 1 wo doses General Safety Test. 1 wo doses microspheres were suspended by sonication for 5 minutes in 3.1 mls of sterile microspheres were suspended by sonication for 5 minutes in 3.1 mls of sterile microspheres were suspended by sonication for 5 minutes in 3.1 mls of sterile
13		microspheres were selected and
		vaccine dilutent consisting of injectable saline containing 0.5% Polysorbate 60°
14		i in medioneally illustration
		vaccine dilutent consisting of injectation and 3 mls N.F., 0.03 ml were injected intraperitoneally into each of 2 mice and 3 mls were administered by gastric lavage to each of 2 guinea pigs. The animals were administered by gastric lavage to each of 2 guinea pigs.
15		· laurage to Each UI -
16		were administered by gassing the vaccine administration.
.0		here and at 7 days following
17		All animals were observed daily for any signs of toxicity. All animals were observed daily for any signs of toxicity.
		All onimals were observed daily for any signs of
18		male specially pure
		Rabbits. 1.5-2 kilogram made a
19)	colony maintained at
20	`	white rabbits, obtained from closed constructions white rabbits, obtained from closed constructions of the study if they did not have of Health, Bethesda, MD. They were selected for study if they did not have of Health, Bethesda, MD. They were selected for study if they did not have
20	,	- THE WALL STREET
2	1	of Health, Bethesian, Parking to CFA/II antigens by HISA and
_		of Health, Bethesda, MD. They would be the of Health, Bethesda, MD. They would be considered by Culture of rectal swales.
2	22	measurable serum antibodies at 1.2 determined by culture of rectal swals. were not colonized by E. coli as determined by culture of rectal swals.
		were not colonized by E. coll as detailed.
7	23	MOIA 1

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	Intra-Muscular Immunization of Rabbits and ELISA. Two Rabbits
	Intra-Muscular Infinity in the phone vaccine at 25 ug protein in two
	Intra-Muscular Immunization were immunized with CFA/II microsphere vaccine at 25 ug protein in two different sites intra-muscularly on day 0. Sera were obtained from all animals
3	different sites intra-muscularly on day 0. Seed 14. The sera were tested by
4	different sites intra-muscularly on the different sites in the diff
5	before immunization on day of and days before immunization on day of and days ELISA for IgG antibodies to CFA/II antigen and individual coli surface (CS) ELISA for IgG antibodies to CFA/II antigen and individual coli surface (CS)
6	Proteins CS3 and CS1. ELISA plates were coated with 3 ug/ml of either proteins CS3 and CS1.
7	proteins CS3 and CS1. ELISA place of the protein (150 ul/well) and incubated with 150 CFA/II antigen, CS3 or CS1 protein (150 ul/well) and incubated with 150
8	DCA for four nouis at 100
9 ,	ul/well of PBS with 0.1% BSA for foot with PBS and 100 ul/well of different dilutions with 0.1% BSA is washed out with PBS and 100 ul/well of different dilutions with 0.1% BSA is washed out with PBS and 100 ul/well of different dilutions ranged
10	with 0.1% BSA is washed out with 120 of each rabbit serum in triplicate was added to the plates. The dilutions ranged of each rabbit serum in triplicate was added to the plates. The dilutions ranged
11	of each rabbit serum in triplicate was a from undiluted to 1:1,000,00. The plates were incubated with the sera for 3
12	from undiluted to 1:1,000,00. The photos and the horse radish hours at 37° C. The sera were washed out with PBS and then horse radish hours at 37° C.
13	hours at 37° C. The sera were washed to the plates at a 1:1000 peroxidase-conjugated goat anti-rabbit IgG was added to the plates at a 1:1000
14	ma plates were incubated to
15	acceptions conjugate. The conjugate
16	- 4 100 ul/well of an ABIS succession
17	was added to care.
18	plates were read using the ELISA reader(DynateLit Editors)
19	of 405 nm after 15 minutes.
20	and expressed as andoocy and
	a straightion of Rabbilla Rabbilla
21	containing either 23 of 30 and
22	and certaining 0.5% Polyson
23	suspended in 1 ml of PBS containing some suspended in 1 ml of PBS containing s
24	sonication. The history

bronchoscope into the duodenum of the rabbits following sedation with an intra muscular injection of ketamine HCl (50 mgm I.M.)(Ketaset, Fort Dodge Laboratories, Fort Dodge, IA) and Lylazine (10 mgm I.M.) (Rompom Malay Corporation, Shnanee, KS). The endoscope was advanced ready under direct vission into the stomach which was insufflated with a 50 ml bolus of room air via a catheter passed through the biospy channel. The catheter was advanced through the pylorus 3-4 cm into the duodemum and the microsphere suspension in 1 ml of PBS was injected, followed by a 9 ml flush of PBS and removal of the air bolus. The rabbits were sacrified by chemical euthanasia at day 14.

Anti-CFA/II Stimulated Lymphocyted Transformation. The Peyer's Patchs were removed and cell suspension obtained by teasing and irigation with a 20 guage needle and syringe. The cells were placed in 2 ml of media at

Patchs were removed and cell suspension obtained by teasing and irigation with a 20 guage needle and syringe. The cells were placed in 2 ml of media at a concentration of 2.5 x 10° cells/ml for each well of a 24 well plate. These cells were challenged separately with BSA and the CFA/II antigen at doses of 500, 50 and 5 ng/ml in triplicate wells. The plates were incubated at 37° C with 5% CO₃. On day 4 the cells were mixed while still inside the wells and 100 ul were transferred into each of 4 wells in a 96 well flat bottom microculture plate. Thus, the challenge at each antigen dose represented by 3 wells in the 24 well plate is now represented by 12 wells in the 96 wdl plate. After the cells have been transferred, each well is pulsed with 20ul of 50 uCi/ml tritiated thymidine. These pulsed plates were incubated for 6 km them harvester with 96 Mach II Cell harvested (Tourtec, Inc.). The lymphocyte proliferation was determined by the tritriated thymidine incoporation measured in kilo counts per minute (Kcpm) using the 1205 Beta Plate Liquid scanification

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counter (LKB, Wallac, Inc.). The results are expressed as mean Kepm + SD and compared to media controls.

Anti-CFA/II Antibody Secreting B Cells. Spleen cells were obtained from immunized rabbits on day 14 following intra-duodenal immunization with CFA/II microsphere vaccine. The cells were placed in 96 well round bottom microculture plate at a final concentration of 6 x 10° cells/well and incubated for 0, 1, 2, 3, 4 and 5 days at 37 C with 5 CO₂. 96 well flat bottom microculture plates were coated with 3 ug/ml of CFA/II antigen overnight blocked with PBS with 0.05% Polysorbate 60°. On the harvest days, the cells were gently flushed out of the wells of the round bottom plates and transferred to the corresponding well in the antigen coated, 96 well flat bottom microculture plates to be tested for the presence of antibody secreting cells using ELISPOT technique. The plates were incubated with the cells overnight at 4° C. The cells were then washed out of the flat bottom plates with PBS, and 100 ul/well of horserudish-peroxidase conjugated, goat anti-rabbit total antibody (IgM, IgG, and IgA) at a 1:1000 dilution were added to the plates. The Plates were incubated for 1 hour at room temperature, at which time, the conjugate was washed out of the plates with PBS. 0.1 mgm of agamse was dissolved in 10 ml of PBS by boiling. After the agar solution cooled but and hardened, 6 mgm of 4-chloro-naphthol, 2 mls of methanol and 30 ulof hydrogen peroxide were added to make the substrate solution. The solution was placed into the flat bottom plates (100 ul/well) and the plates were held at 4. C overnight so the agar could harden. The number of browish spec 15

wells (total of 9 x 10° spleen cells) was counted and represents the number of 1 antibody secreting cells per 9 x 10° spleen cells. Pathological Evaluation. Rabbits were euthanized by parenteral 2 overdose of sodium pentobarbital and were subjected to complete 3 4 necropsy. Sample of tissue including small and large intestine with gut associated lymphoid tissue, spleen, mesenteric and mediastinal lymph 5 nodes, lung, trachea, liver and kidney were fixed by immersion in 10% neutral 6 buffered formalin. Tissues were routinely processed for light microscopy and 7 embedded in paraffin. Five micron thick sections were stained with 8 9 hematoxylin and eosin. Statistical Analysis. The paired student t-test was used to determine p 10 11 values. 12 RESULTS Particle Size Distribution. The results of size frequency analysis of 13 150 randomly chosen microspheres are shown in (Figure 37). The particle 14 size distribution is plotted in % frequency against particle size in diameter 15 (size) expressed in um. The average number fregency diameter is 4.8 um. 16 The average volume frequency diameter is 4.6 um. The percent volume 17 between diameters of 5-10 um is 63% and the percent volume less than 10mm 18 19 diameter is 88%. Scanning Electoron Microscopy. The microspheres are seen in 20 (Figure 38) which is a scanning electron photomicrograph. Nearly all the 21 22

•	microspheres are less than 10 um as compared to the 5 um bar. Also the
1	surfaces of the microsphere are smooth and demonstrate lack of pores.
2	Protein Content. The protein loads of the individual batches are the
3	Protein Content. The protein value of the protein Content. The protein Content. The protein Content. The protein Content.
4	following: K62A8, 1.16% ± 0.10 SD; K63A8, 1.023% ± 0.17SD; K64A8,
5	1.232% ± 0.13 SD; and K65A8, 0.966% ± 0.128 SD. The mean
6	The protein load is 1.16% ± 0.15 SD. The protein load of the CFA/H
7	microsphere vaccine in the final dose vial is the following: Lot L74F2,
8	1.175% ± 0.17SD.
	Moissure Content. The CFA/II microsphere vaccine (Lot /4P2)
9	percent water content was found using the Karl Fischer titrimeter method to be
11	a 1547 using triplicate samples.
	A seconitrile and Heptane Residuals. The acetonitrile residuals of the 4
12	individual CFA/II microsphere batches are the following: K62A8, <0.1%;
13	individual CFA/11 interespective
14	K62A8, <0.1%, K64A8, <0.1%, min the final dose vial is the
15	residual of the CFA/II microsphere vaccine in the final dose vial is the
16	following: Lot L74F2, 0.07 ± 0.05%. The heptane residual of the 4
17	CEA/II microsphere batches are the following: K62A8, 1.3A,
18	Following pooling to
	to access and subsequent drying, the heptane residual of the CFA/II microspace
19	to the final dose vial is the following: Lot L74F2, 1.6 ± 0.13.
20	Microbial load. One hundred milligrams (a single dose) of CFA/II
21	microsphere vaccine (Lot L74F2) in the final dose vial was suspended in 2 2
22	microsphere vaccine (Lox L/4/2) in and a blood again culture plate x 2.
23	microsphere vaccard 1 ml poured onto a blood agar culture plate x 2. ml of sterile saline and 1 ml poured onto a blood agar culture plate x 2.
24	Twenty two colonies grew after 48 hours of culture and 21 were identified as

bacteria are considered to be nonpathogenic to humans. An additional 100 mgms of CFA/II microsphere vaccine (Lot L74F2) were suspended in 2 ml of sterile saline and 0.25 ml poured onto four different fungal culture agars and cultered for 5 weeks. Three fungal colonies grew and each was identified as A. glaucus.

Were incubated each in 1 ml of PBS, pH 7.4 at 37° C for 0, 1, 3, 6, 8, 15 and 22 hours. The superanates were removed and replaced at these times. The protein content was determined for each supernate sample and the results are seen in (Figure # 39). The results are plotted as percent release of CFA/II against time in hours. An average of 8% of CFA/II is released at one hour rising to 20% at 8 hours then a slower release to 25% at 22 hours.

General Safety Test. Two one hundred milligrams(a single dose) of CFA/II microsphere vaccine in the final dose vials were suspended in 3.1 mls of the sterile dilulent consisting of 0.85 N saline prepared for injection plus Polysorbrate 60° at 0.5%. Two Swiss mice (16.5 gm) were injected intraperitoneally with 0.03 mls and two Hartley guinea pigs (350 gm) were administered by gastric lavage 3.0 mls.

None of these animals displayed any signs of toxicity for 7 days. The mice gained and average of 2.3 gms and the guinea pigs gained and average, of 43 grams. The CFA/II microsphere vacccine therefore passed the general safety test.

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	Serum IgG Antibody Responses. Two rabbits were immunized in two
	separate sites intra-muscularly with 25 ug of protein of CFA/II microsphere
	separate sites intra-muscularly with 200 separate sites in 200 separate sites in 200 separate sites sites in 200 separate sites sites in 200 separate sites sit
	vaccine (Lot L74F2) in the final dose vial. Sera samples were obtained before
	and 7 and 14 days following immunization. The IgG antibody titers to CFA/II
	and the results were determined using ELISA and the results were in
	(Figure 32). The results are expressed as mean antibody titers against the
	(Figure 32). The results are expressed an involve titers greater than 1000
	different antigens at 0, 7 and 14 days. High antibody titers greater than 1000
	were seen at 7 days to both CS1 and CS3 protein which rose to greater than
1	is an account day 14. The individuals titers to CFA/II are seen in (Figure 33).
)	Rabbit 109 developed an antibody titer of 1,000 by day 7 rising to 3,000 by
)	Rabbit 109 developed an antibody developed and dev
1	day 14. Rabbit 108 had a log higher rise at day 7 and 2 log higher rise at day
2	14 being 3 x 10° at day 7 going to 1 x 10° at day 14.
	A: CEA/II Stimulated Lymphocyte Transformation. Pive rapping
3	were immunized intra-duodenally with CFA/II microspheres containing either
4	were immunized intra-duodentary
15	were immunized into a coordinate into a coordina
16	and then sacrificied on day 14. The Peyer's patch lymphocytes were
	challenged in vitro with CFA/II antigen, BSA media and alone. The
17	lymphocyte transformation was determined by tritriated thymidine
18	lymphocyte transformation was description are seen in (Figure
19	lymphocyte transformation will be light dose immunization are seen in (Figure incorporation. The results of the high dose immunization are seen in (Figure
	The agree expressed as Kepm against antigen dose. Two respects to
20	and a media control is seen in any of the five rabbits. All rabbits responses
21	by lymphocyte transformation in a dose dependent manner to the CFAII.
22	by lymphocyte transformation in a control are highly. The highest dose responses were 3-10X's the media control are highly
23	The highest dose responses were 3-10-10 to the first receiving
	significant with a p value of <0.002. The results of the 5 rabbits receiving
24	

the low dose immunization are seen in (Figures 35). Rabbit #80 gave no response probably due to poor Peyer's patch cell population which did not respond were to Conconavallin A mitogenic stimulation either. The remaining 4 rabbits gave positive responses with the high CFA/II dose response being 2-8x media control and highly significant with p values of <0.009. Again no response were seen to BSA compared to the media cont

intraduodenally with CFA/II microsphere containing 50 ug of CFA/II protein at days 0, 7 than sacrified at day 14 were studied. The spleen cells were placed into microculture then ELISPOT forming B-Cells secreting specific anti CFA/II antibody determined at days 0, 1, 2, 3, 4 and 5. The results are seen in (Figure 36) and expressed as # of antibody secreting cells per 9 x 10 spleen cell against culture days. Positive responses were seen in all 5 rabbits on days 2-5. Days of maximum responses occurred on day 3 for rabbits 65 and 66; day 4 for rabbit 85; amd day 5 for rabbits 83 and 86. The responses are highly significant being 7-115 times higher than the 1-2 cells seen on all days in 4 control rabbit (67, 69, 72, 89) (Figure 45). Here is a composite graph expressing the mean counts ± ISD for all days of culture.

Pathological Evaluation. A consistent finding in the spleens of all rabbits both the 25 and 50 ug protein dose groups was minimal to mid diffuse lymphocytic hyperplasia the periarteriolar lymphatic sheaths (T cell dependent areas). Two of five rabbits of the 50 ug dose group (#83 and #86) also had mild lymphocytic hyperplasia of splenic follicular (B cell dependent) areas.

The three rabbits in an untreated control group had histologically normal 1 spleens. 2 Reactive hyperplasia of mesenteric lymph nodes was often seen in 3 vaccinated rabbits. Two of five rabbits in the 25 ug dose equivalent group (#83 and #86) also had minimal to mild lymphocytic hyperplasia of cortical 4 follicular (B cell dependent) areas. The mesenteric lymph nodes of the other 5 6 vaccinated rabbits and of the untreated control rabbits were within normal 7 limits. Incidental or background lesions found in one or more rabbits of all 8 three group were acute minimal to mild pnuemonia and foreign body 9 microgranulomas of the cecal gut associated lymphoid tissue. 10 Disscussion McQueen et al (33)has found that the AF/R1 adhesin of rabbit diarrheagenic 11 Escherichai coli (RDEC-1) incorporated into biodegrable microspheres could 12 13 function as a safe and effective oral intestinal vaccine in the rabbit diarrhea 14 model. The AF/R1 was incorporated into poly D,L-lactide-co-glycolide) microspheres and administered intraduodenally. Jarboe et al (34) reponed that 15 16 Peyer's patch cells obtained from rabbits immunized intra-duodenually 17 with AF/R1 in microspheres responded with lymphocte proliferation spea in 18 vitro challenge with AF/R1. This early response at 14 days gave a char 19 indication as to the immunogenicity of E. coli pili contained within the 20 polymer microspheres. 21 In developing an effective oral vaccine against enterotoxigen's E. odi. CFA/II pili given as an oral vaccine was found to be ineffective. The CFA/II 22 pilus proteins were found to be rapidly degraded when treated with Olaiki 23

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and pepsin conditions mimicking those contained in the stomach (27). The CFA/II was found to be immunogenic when given in high doses intraintestinally producing intestinal secretary IgA antibodies (26).

The CFA/II vaccine has now been incorporated into poly(D,L lactideco-glycolide) microspheres under Good Manufacturing Practices and tested under Good Laboratory Practices. The microspheres, are spherical, smooth surfaced and without pores. The majority (63%) are between 5-10 um in diameter by volume. This size range has been suggested to promote localization within the Peyer's patch in mice and perhaps enhance local immunization (29-32). The protein content being 1.174% is close to 1% which was the goal of the vaccine formulation. One percent was chosen because 0.62% was the core loading of the AF/R1 microspheres which were effective. Also a small precentage perhaps 1-5% (35) is anticipated to be taken up from the intestine, a higher protein content would lead to considerable loss of protein.

The organic residuals are of course a concern. Heptane exposure would be 1.7 mgm per vaccine dose. This is compared to the occupational maximum allowable exposure of 1800 mgm/15 min. Therefore, the heptane contained with the CFA/II microsphere vaccine appears to be a safe level. The acetonitrile is very low - 0.1 mgm per vaccine dose. The human oral TDLO is 570 mgm/Kg (any non letheal toxicity). Therefore, the accuminitie contained with the CFA/II microsphere vaccine appears to be at a safe level. The CFA/II vaccine was produced under sterile conditions. However, the process of incorporation of the desalted CFA/II vaccine into the polymer

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The antibody secreting B-cells demonstrated in the rabbit spleen at 14
days is a clear indication that B-cells have been immunized. They may
represent resident B-cells immunized in the spleen or B-cells immunized at the
level of the Peyer's patches and are migrating through the spleen to return to
the intestial mucosal lamina propria (1-3). The delay of several days before
secreted antibody is detected suggests either manuration is required of the B-
cells or that down regulation may be present initially and lost with time in
culture.
Further evidence of immunization by the CFA/II microsphere vaccine
given intra-duodenually is demonstrated by the lymphatic hyperplasia in the
spleen seen to a greater extend in the rabbits receiving the lower dose 5/5
compared to 2/5 of the rabbits receiving the higher 50 ug protein dose. On the
other hand, greater T-cell dependent area lymphoytic hyperplasia in the
mesenteric lymph nodes were seen in rabbits receiving the higher 50 ug dose,
4/5 compared to 2/5. These changes are most likely due to the vaccine since

abnormal pathological changes attributable to the vaccine were seen.

The CFA/II BPM vaccine has undergone pre-clinical evaluation and has been found safe and immunogenic. This vaccine is ready for clinical Part I safety testing following FDA's IND approval.

similar changes were not seen in three untreated control rabbits. Also no

PART : III

In sum, alum precipitation, vaccination regimen and controlled delivery by microencapsulation were studied to determine what criteria must be satisfied to provide a protective immune response to hepatitis B surface antigen.

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(HBsAg) after a single injection of vaccine. In mouse studies, the 50% effective dose (EDs) for the alum precipitated Heptavax B vaccine (Merck, Sharp and Dohme) was 3.8 ng when administered in a 3 injection regimen, but was 130 ng when one immunizing dose was used. Antigen release studies revealed that HBsAg is bound tightly to the alum, indicating that the antigen remains in situ until scavenged by phagocytic cells. the ED₅₀ with a 3 dose regimen of aqueous HBsAg was 180 ng, a opposed to over 2000 ng for daily injections of low doses for 90 days and 240 ng for a regimen that employed initially high doses that decreased geometrically at 3 day intervals over 90 days. The ED₅₀ was 220 ng for a single dose regimen of HBsAg microencapsulated in poly (DL-lactide-co-glycolide) in a form that was too large to be phagocytized and had an antigen release profile similar to that achieved with the geometrically decreasing regimen of doses. This indicates that single injection of microencapsulated immunogens can achieve similar effects in vivo to those achieved with multiple dose regimens. For HBsAg the effect to be achieved appears to be 3 pulses of particulate immunogens that can be scavenged by phagocytes.

INTRODUCTION

A major disadvantage of inactivated vaccines lies in their inability to confer lasting immunity. Due to rapid elimination from the body, multiple doses and boosters are usually required for continued protection³⁷. Alum adjuvants, achieving their effects by mechanisms of antigen presentation and

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sustained antigen release³⁴, have been used successfully to increase the potency of several inactivated vaccines including those against tetanus, anthrax, and serum hepatitis39,40. Though useful, alum preparations are deficient in several aspects. Control over quantity and rate of antigen release is limited, often resulting in a continued requirement for immunization schedules consisting of multiple injections given over a period of several months to years. Alum adjuvants are also non-biodegradable and thus remain within the body, serving as a nidus for scar tissue formation38 long after they have served their function.

Protracted, multiple immunization schedules are unacceptable during massive mobilization and deployment of troops. Changing global disease patterns and deployment of new biological warfare agents by enemy forces require flexibility in the number and types of vaccine antigen administered to soldiers departing for combat. Any immunization schedule requiring completion during engagement in non-linear combat would compromise this flexibility and place an unreasonable burden on our health care delivery system.

The main objective of this study was, therefore, to develop a biodegradable, controlled-release adjuvant system capable of eliminating the need for multistep vaccination schedules. This investigation was designed to: (1) determine in an animal model hepatitis B vaccine release rate characteristics desirable for single-step immunization, (2) incorporate those release rate characteristics into a one-step biodegradable poly(DL-lacide-coglycolide)(DL-PLG) microencapsulated hepatitis B surface antigen (HBsAg) vaccine, and (3) conduct an in vivo trial comparing the effectiveness of this

WO 98/32427 PCT/US98/01556

	single-step vaccine against the conventional three-step hepatitis vaccine
1	currently employed. The results were intended to provide the foundation for
2	currently employed. The results were against hepatitis and other
3	further development of single step vaccines against hepatitis and other
4	militarily significant diseases ¹² .

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MATERIALS AND METHODS

Vaccine potency assay. Due to its availability, compatibility with cage mates, and potential application to the study of hepatitis B vaccine¹³, the female Walter Reed (ICR) stain mouse was used. A hepatitis B vaccine potency assay for comparing the six-month immunization schedule currently in use⁴¹ with that of a single-step immunization by sustained antigen release was established according to the following protocol: Specimens for baseline antibody titers were collected from twenty mice by exsanguination. Immediately prior to exsanguination, all mice employed in this and other exsanguination procedures in these studies were anesthetized with a 0.1 ml injection of V-Pento. Groups of 12 mice were then immunized according to a schedule consisting of either 0.25 ug, 0.025 ug, 2.5 ng, 0.25 ng, 2.5 pg, or 0.25 pg Heptavax B vaccine (HBV) administered in 50 microliter volumes subcutaneously (s.c.) at the beginning and end of the first, and end of the second month of the protocol. Antibody responses to the vaccine were monitored immediately before the third injection and approximately one month after the third injection. Specimens for antibody determination were collected by exsanguination of seven anesthetized mice from each group and assayed along with the baseline samples by the Abbott Ausab radioimmunoassay. Percent seroconversion verses micrograms vaccine employed with calculated by the method of Reed and Muench⁴³. These data were employed to establish a mouse vaccine potency assay calibrated to detect differences between Heptavax B and other forms of hepatitis b vaccine.

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In vitro antigen release rate from Heptavax B vaccine. Antigen release from aluminum hydroxide adjuvant in HBV was measured by pumping 2 cc per hour of 1:20,000 thimerosal in saline at 4°c across a 0.2 u pore diameter Acrodisc filter apparatus containing 20 ug of vaccine. The effluent, collected by a Gilford fraction collector, was assayed periodically over several weeks for protein by UV absorption at 280 nm on a Beckman model 25 double beam spectrophotometer, and for HBsAg by the Abbot Ausria II radioimmunoassay made quantitative by using HBsAg standards supplied by Merk, Sharp, and Dohme. Accuracy of the HBsAg standards were verified by Biuret protein determination and by UV absorbance at 215 nm and 225 nm⁴⁴. Nonspecific antigen retention on the Acrodisc filter was assessed by measuring percent recovery of a known quantity of HBsAg. Spontaneous degradation of vaccine antigen was monitored by comparing daily rations of antigen to total protein detected in the effluent.

characterize the stability of the aqueous antigen to the various physical conditions employed in the microencapsulation process. Conditions tested included lyophilization with reconstitution in distilled water, cyclohexane, methylene chloride, chloroform, methyl alcohol, acetone, iso-octane, hexane, acetone, pentane, or heptane; irradiation while lyophilized; and, exposure to elevated temperatures. Samples exposed to organic solvents were first lyophilized, reconstituted with the test solvent, evaporated to dryness under nitrogen at room temperature and reconstituted with distilled water. Test samples were compared against untreated controls by assaying serial dilutions

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	of each with the Abbot Ausria II procedure and comparing the plots of counts
1	per minute verses dilution.
2	Assessment of the effect of antigen release rate on vaccine potency.
3	Three regimens simulating patterns of free HBsAg release that could be
4	Three regimens simulating patterns of the same and with the three monthly dose
5	achieved by microencapsulation were contrasted with the three monthly dose
6	regimen of Heptavax B for immunizing mice. To do so, 24 ICR mice were
7	divided into groups and vaccinated as indicated below. Seven mice from each
8	subgroup were exsanguinated at the end of the second and third months of the
9	experiment. The sera were separated and assayed for specific antibody
	response to HBsAg by Abbot Ausab procedure.
10	HV regimen a: 14 mice/treatment receiving 3 s.c. injections of 250,
11	25, 2.5 or 0.25 ng doses of HBV a month apart.
12	HBsAg regimen a: 14 mice/treatment receiving 3 s.c. injections of
13	250, 25, 2.5 or 0.25 ng doses of aqueous HBsAg a month apart.
14	250, 25, 2.5 or 0.25 ng doses of aqueent receiving total doses of 750, 75, HBsAg regimen b: 14 mice/treatment receiving total doses of 750, 75,
15	HBsAg regimen b: 14 mice/treatment receiving the HBsAg regimen receiving the HBsAg receiving the HBs
16	7.5 or 0.75 ng of aqueous HBsAg over 3 months by s.c. injections of ZX _Y ng
17	at 3 day intervals, where Z is the total dose, y is the injection number, and X
18	is the fraction indicated on the graph in Fig. 1 minus the fraction for the
19	vious injection.
	HBsAg regimen c: 14 mice/treatment receiving daily s.c. injections of
20	8.33, 0.833, 0.0833 or 0.00833 ng of aqueous HBsAg for 3 months.
21	Microencapsulation in DL:PLG. Microencapsulated immunogens
22	were fabricated by Southern Research Institute, Birmingham, AL. DLPIG
23	were fabricated by Southern Research Library, DL lactide and glycolide, polymers were synthesized from the cyclic diesters, DL lactide and glycolide,
24	polymers were synthesized from the cyclic discussion

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by using a ring-opening melt polymerization catalyzed by tetraphenyl tines. The resulting polymer was dissolved i methylene chloride, filtered free of insoluble contaminants and precipitated in methanol. Lactide-co-glycolide mole ration of the product was determined by nuclear magnetic resonance spectroscopy. Encapsulation of HBsAg in DL:PLG polymer was achieved by an organic phase separation process46. Microcapsules of the desired size (approximately 100 micron diameter in these studies) were isolated from each batch by wet sieving with hexane through standard mesh stainless steel sieves and then dried for 24 hours in a vacuum chamber maintained at room temperature.

In vitro analysis of encapsulated antigens. Integrity of encapsulated antigen was assessed by comparing the antigen to total protein ratios present in microcapsule hydrolysates with those obtained from suspensions of pure unencapsulated antigen. Centrifuge tubes containing 1 ug of either microencapsulated or pure vaccine antigen in 1 ml saline were incubated at 4,c with shaking. Samples were collected at weekly intervals by interrupting the incubation, sedimenting the contents of the tubes by centrifugation and withdrawing the supernates. Sediments were resuspended in 200 microliters of saline and supernates were assayed for HBsAg by the Abbott Ausria II radioimmunoassay. The HBsAg standard described earlier in this report was used as the calibrator. Antigen destruction due to the encapsulation procedure was monitored by a comparison between the antigen assayed from the hydrolysate and from the untreated antigen control.

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	Assessment of the potency of DL:PLG microencapsulated HBsAg for
	immunizing ICR mice when used alone and in combination with Heptavax B
	and a leaded microcapsules that had been rabricated by bottom
	Research Institute to release the majority of their HBsAg load within 40 to 50
	Research Institute to release the majorny
	Research Institute to Ichean and Institute to Ichean and Institute to Ichean and Ichean
•	and composition. The research
)	microcapsule preparations were placed onto lyophia.
7	not in use in order to assure minimum spontaneous degradation prior to
8	not in use in order to assure minimum injection. On the day of injection, a predetermined weight of microcapsules or injection.
9	injection. On the day of injection, a predetermine
0	placebo-diluted microcapsules was added to each syringe. Immediately prior
_	of injection vehicle (2 Wt % Carbox).
11	Tween 240 in water, Southern Research Historical water
12	drawn into the microcapsule-loaded syringes, mixed and injected. All mice
13	drawn into the microcapsule-loaded symmetry
14	were vaccinated s.c. as indicated below:
	Group 1: 14 mice/treatment receiving 25, 25, 2.5, 0.25 or 0.925 ng
15	
16	HBV. Group 2: 14 mice/treatment receiving 1000, 250, 25 or 2.5 ng
17	Group 2: 14 mice dealines (BSA)
18	aqueous HBsAg with Bovine Serum Albumin (BSA).
	Group 3: 7 mice receiving 1600 ng microencapsulated HBsAg
19	(HBsAg) plus 0.25 ng HBV and 14 mice/treatment receiving 160, 16, 1.6 or
20	
21	0.16 ng HBsAg plus 0.25 ng HBV. Group 4: 7 mice receiving 1600 ng HBsAg plus 2.5 ng HBV and 14
22	Group 4: 7 mice receiving 1000 ing 12 Track a plus 2.5 ng HBV.
23	mice/treatment receiving 160, 16, 1.6 or 0.16 ng HBsAg plus 2.5 ng HBV.

1	Group 5.: 7 mice receiving 1600 ng HBsAg plus 25 ng HBV and 14
2	mice/treatment receiving 160, 16, 1.6 or 0.16 ng HBsAg plus 25 ng NBV.
	Group 6: 7 mice receiving 2500 ng HBsAg and 14 mice-treatment
3	receiving 250, 25, 2.5 or 0.25 ng HBsAg. Fifty-three days after receiving the
4	above injections, the mice were anesthetized with an 0.1 cc injection of V-
5	Pento and exsanguinated. Blood samples were allowed clot and the sera were
6	separated by centrifugation. The serum samples were assayed for antibody to
7.	HBsAg by the Abbott Ausab procedure.
8	RESULTS
9	Heptavax B vaccine potency. As can be seen from Table 4, the total
10	
11	dose of vaccine which produced seroconversion in 50% of
	- ICP mice
12	TABLE 1.2 Potency of Heptavax B vaccine in ICR mice.
13	ED.
14	No. ng Heptavax B per Injection ED50
15	Inj. 250 25 2.5 .25 .025 .0025 .00025 ng
16	
17	2 5/5 4/4 3/6 2/6 0/5 1/4 0/4 1.7
18	3 6/6 6/6 4/6 1/6 0/6 1/6 1/6 2.0
19	
20	 Number positive seroconversions per number vaccinated.
	The vaccinated mice (ED ₅₀) for HBV was approximately 2 ng, whether the
21	vaccine was given in 2 or 3 injections.
22	Addition and D

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In vitro antigen release rate from HBV. HBsAg release from the 20 ug of Heptavax was not detected in any of the 21 fractions of saline collected from the Acrodisc polycarbonate filter over a 30 day period. The lower limit of detection for the Abbott Auria II assay employed was approximately 4.8 ng/ml. The Acrodisc filter used in the antigen release study was back-washed with 10 mls normal saline. Quantitation of the HBsAg present within this back-wash eluent revealed the presence of the original 40 ug of Heptavax vaccine which had been loaded into the filter at the start of the experiment. - This is the concentration which one would expect to obtain if there had been no deterioration of the original 40 ug/ml HBsAg loaded onto the filter, none of the antigen eluted from the alum adjuvant, and none of the vaccine had adsorbed onto or passed through the filter.

Evaluation of antigen stability. Considerable effort was expended in assessing the effects of physical conditions on the antigenicity of HBsAg to insure that the conditions used for microencapsulation would not cause serious degradation of the immunogen. Since microencapsulation must be performed on dried materials which are suspended in organic solvents, the HBsAg, which was provided as a solution, had to be lyophilized. Initial attempts at lyophilizing HBsAg in normal saline resulted in a total loss of detectable antigen within samples. Dilution of the HBsAg sample 1:10 in distilled water prior to freezing resulted in reservation of nearly 100% of the antigen detectable in the original sample. Studies of antigen stability at elevated temperature revealed that HBsAg may be heated to 50°c for up to one hour without appreciable loss of antigen. The studies involving exposure of

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lyophilized antigen to organic solvents indicated that iso-cane and nexale had
minimal effects on antigenicity, but that 95% to 100% of antigenicity was lost
upon exposure to either methylene chloride, chloroform, cyclohexane, or
methyl alcohol. Moderate antigen loss occurred in the presence of acetone,
pentane and heptane. As a result of these studies, hexane was chosen as the
solvent for microencapsulation.
Assessment of the effect of antigen release rate on vaccine potency.
The results (Tablel 3) indicated that immunogen formation (i.e., the alum
adjuvant of Heptavax B) had far more

1	TABLE 13 Effect of immunogen formulation and vaccination
2	regimen on potency for immunizing ICR mice.
3	
4	Immunogen ng Total Dose HBsAg ED ₅₀
5	Formulation Regiment 750 75 7.5 .75 ng
6	
7	Heptavax B a 7/7° 6/6 5/7 1/7 3.8
8	Aqu. HBsAg a 4/6 3/7 0/7 0/6 180
9	Aqu. HBsAg b 6/7 0/7 1/7 0/7 240
10	Aqu. HBsAg c 1/7 0/7 0/7 > 2000
11	
12	 Number positive seroconversions per number vaccinated.
13	a 3 injections of 1/3 total dose a month apart.
14	b Injections administered every three days for 90 days in
15	decreasing dosages according to a logarithmic progression.
16	c Injections of 1/90 total dose daily for 90 days.
	the and that pulsing with large
17	effect on potency than did the vaccination regimen, and that pulsing with large
18	doses of immunogen was more effective than continuous administration of
19	small doses.
20	HBsAg release from DL:PLG microcapsules. The microcapsules
21	employed in this study were designed to disintegrate within three weeks after
22	hydration. It is evident from the release curve (Fig.10) that they performed as
23	designed, releasing approximately 17% of their total load in an initial pulse

and approximately 7% of the remaining available HBsAg over the first three weeks. Assessment of the potency of DL:PLG microencapsulated HBsAg for immunizing ICR mice when used alone and in combination with Heptavax B vaccine. The results (Table 14) indicate that the microencapsulated HBsAg ha approximately the same immunogenicity as did the Heptavax B. Neither immunogens were sufficiently potent to effect with a singly injection seroconversion rates similar to those achieved after three injections of Heptavax B (Table 12). Only the immunogen TABLE 14 Potencies of Heptavax B and microencapsulated HBsAg by single injection S.C. who administered alone and in combination to immunize ICR mice. Var. Dose ng Coast. ng Variable Dose Var. Dose Tot. Dose lmmunogen Dose mHBsAg 2500 250 25 2.5 .25 ED _m ng ED _m ng Heptavax B 0 13/14* 2/14 4/14 0/13 130 130 Heptavax B 1.6 10/13 1/14 0/13 100 100 Heptavax B 1.6 10/13 1/14 0/13 100 100 Heptavax B 160 3/12 2/11 1/12 >370 >530 Heptavax B 160 7/7 7/7 7/7 <0.8 1600 Mic. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220		
Assessment of the potency of DL:PLG microencapsulated HBsAg for immunizing ICR mice when used alone and in combination with Heptavax B vaccine. The results (Table 1) indicate that the microencapsulated HBsAg has approximately the same immunogenicity as did the Heptavax B. Neither immunogens were sufficiently potent to effect with a singly injection seroconversion rates similar to those achieved after three injections of Heptavax B (Table 1). Only the immunogen TABLE 1/4 Potencies of Heptavax B and microencapsulated HBsAg by single injection S.C. who administered alone and in combination to immunize ICR mice. Var. Dose ng Const. ng Variable Dose Var. Dose Tot. Dose Immunogen Dose mHBsAg 2500 250 25 2.5 .25 ED _m ng ED _m ng Heptavax B 0 13/14* 8/14 4/14 0/13 130 130 Heptavax B 0.16 11/13 4/14 1/14 1.7 1.8 Heptavax B 1.6 10/13 1/14 0/13. 100 100 Heptavax B 160 3/14 1/14 1/14 > 470 > 490 Heptavax B 160 3/12 2/11 1/12 > 370 > 530 Heptavax B 1600 7/7 7/7 7/7 < 0.8 1600 MGe. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220	1	and approximately 7% of the remaining available HBsAg over the first three
immunizing ICR mice when used alone and in combination with Heptavax B vaccine. The results (Table 1) indicate that the microencapsulated HBsAg ha approximately the same immunogenicity as did the Heptavax B. Neither immunogens were sufficiently potent to effect with a singly injection seroconversion rates similar to those achieved after three injections of Heptavax B (Table 1). Only the immunogen TABLE 1/4 Potencies of Heptavax B and microencapsulated HBsAg by single injection S.C. wh administered alone and in combination to immunize ICR mice. Var. Dose ng Const. ng Variable Dose Var. Dose Tot. Dose Immunogen Dose mHBsAg 2500 250 25 2.5 .25 EDm ng EDm ng Heptavax B 0 13/14° 8/14 4/14 0/13 130 130 Heptavax B 0.16 11/13 4/14 1/14 1.7 1.8 Heptavax B 1.6 10/13 1/14 0/13 100 100 Heptavax B 16 3/14 1/14 1/14 > 470 > 490 Heptavax B 160 3/12 2/11 1/12 > 370 > 530 Heptavax B 1600 7/7 7/7 7/7 < 0.8 1600 Mie. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220	2	.weeks.
immunizing ICR mice when used alone and in combination with Heptavax B vaccine. The results (Table 1) indicate that the microencapsulated HBsAg ha approximately the same immunogenicity as did the Heptavax B. Neither immunogens were sufficiently potent to effect with a singly injection seroconversion rates similar to those achieved after three injections of Heptavax B (Table 1). Only the immunogen TABLE 1/4 Potencies of Heptavax B and microencapsulated HBsAg by single injection S.C. wh administered alone and in combination to immunize ICR mice. Var. Dose ng Const. ng Variable Dose Var. Dose Tot. Dose Immunogen Dose mHBsAg 2500 250 25 2.5 .25 EDm ng EDm ng Heptavax B 0 13/14° 8/14 4/14 0/13 130 130 Heptavax B 0.16 11/13 4/14 1/14 1.7 1.8 Heptavax B 1.6 10/13 1/14 0/13 100 100 Heptavax B 16 3/14 1/14 1/14 > 470 > 490 Heptavax B 160 3/12 2/11 1/12 > 370 > 530 Heptavax B 1600 7/7 7/7 7/7 < 0.8 1600 Mie. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220	3	Assessment of the potency of DL:PLG microencapsulated HBsAg fo
approximately the same immunogenicity as did the Heptavax B. Neither immunogens were sufficiently potent to effect with a singly injection seroconversion rates similar to those achieved after three injections of Heptavax B (Tablel 2. Only the immunogen TABLE 14 Potencies of Heptavax B and microencapsulated HBsAg by single injection S.C. who administered alone and in combination to immunize ICR mice. Var. Dose ng Const. ng Variable Dose Var. Dose Tot. Dose Immunogen Dose mHBsAg 2500 250 25 2.5 .25 ED _m ng ED _m ng Heptavax B 0 13/14* 8/14 4/14 0/13 130 130 Heptavax B 0.16 11/13 4/14 1/14 1.7 1.8 Heptavax B 1.6 10/13 1/14 0/13 100 100 Heptavax B 160 3/12 2/11 1/12 >370 >530 Heptavax B 160 3/12 2/11 1/12 >370 >530 Heptavax B 160 7/7 7/7 7/7 <0.8 1600 Mie. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220		immunizing ICR mice when used alone and in combination with Heptavax B
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17 immunogens were sufficiently potent to effect with a singly injection 18 seroconversion rates similar to those achieved after three injections of 19 Heptavax B (Tablel 2). Only the immunogen 10 TABLE 1.4 Potencies of Heptavax B and microencapsulated HBsAg by single injection S.C. who 11 administered alone and in combination to immunize ICR mice. 12 Var. Dose ng Const. ng Variable Dose Var. Dose Tot. Dose 14 lmmunogen Dose mHBsAg 2500 250 25 2.5 .25 ED ₂₀ ng ED ₂₀ ng 15 Heptavax B 0 13/14° 8/14 4/14 0/13 130 130 17 Heptavax B 0.16 11/13 4/14 1/14 1.7 1.8 18 Heptavax B 1.6 10/13 1/14 0/13 100 100 19 Heptavax B 16 3/14 1/14 1/14 > 470 > 490 20 Heptavax B 1600 7/7 7/7 7/7 < 0.8 1600 21 Mic. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220	5	vaccing. The rest impurposenicity as did the Heptavax B. Neither
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20 Heptavax B 160 3/12 2/11 1/12 >370 >530 21 Heptavax B 1600 7/7 7/7 7/7 <0.8 1600 22 Mic. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220		Replayan 5 1/14 1/14 >470 >490
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21 Heptavas 3 1000 22 Mie. HBsAg 0 3/6 6/15 1/13 2/10 2/14 220 220		Heptavax B 100
		nepavad 8 1000
		Mic. HBsAg

Number positive seroconversions per number vaccinated.

combination of Heptavax B with 0.16 ng mHGsAg provided this level of seroconversion. At the ED₅₀ endpoint, the 0.16 ng dose of mHGsAg is approximately 10% of the total dose. Similarly, a small amount of Heptavax B appeared to enhance the immunogenicity of the microencapsulated immunogen, although the combination was clearly less immunogenic when the two formulations were present at equivalent concentrations.

DISCUSSION

The potential advantage of microcapsules lies in their ability to be programmed during fabrication into forms that have quite difference release profiles, including slow and steady release, multiple bursts of antigen over a period of time, or combinations of release forms. Sieving allows choice of microcapsule size, and the ability of DL-PLG to sequester antigen from the host's immune system until release occurs enhances control over exposure of the recipient's immune system to antigen over a sustained period of time. These characteristics provided the impetus for these studies as they indicate potential for achieving the effects of a multiple injection regimen by controlling release in vivo after a single injection.

The results of these studies are important for gaining an under standing of the fundamental differences between the manner in which alum and microcapsules interact with the immune system. The antigen release studies showed that alum firmly bound the antigen on its surface, whereas the microcapsules sequestered the antigen load within the interstices of an immunologically inert polymer. Release of antigen from microcapsules was spontaneous and gradual while antigen release from alum wa probably

enzymatically mediated within host macrophages. Alum thus performed at least two useful functions as an adjuvant: by bearing its entire load of antigen upon its surface, it provided a large single exposure of antigen to the host; and, by being readily phagocytized by host macrophages, it served as a means of targeting the antigen to the immune system.

In order for microcapsules to be efficacious as a vaccine delivery system, a means of incorporating the two properties common to alum adjuvant must be devised. These properties, which where discussed above, are targeting antigen to the immune system and delivering the antigen load in a single concentrated pulse at its target. A gradual, sustained release of free antigen, as was achieved with the 100 micron microcapsules used in these studies, could be expected to elicit an immune response similar to that seen with either regimen b or regimen c (Tablel 3), where multiple injections of small doses were employed. In fact, as shown in Tablel 1, the microencapsulated immunogen elicited a response similar to that achieved with regimen b. This is probably due to the fact that the microcapsules release approximately 10% of their antigenic load immediately after injection.

Microcapsules with extended release patterns tend to be large (>10 microns in diameter) and thus fail to be readily phagocytized. In order for the larger microcapsules with prolonged antigen release characteristics to be efficacious, the antigen eventually released from those microcapsules would have be in a form which targeted and concentrated it within the recipient's immune system. This might be effectively achieved by microencapsulation of

	increasing clusters of smaller (< 10
1	antigen coated alum or by microencapsulating clusters of smaller (<10
2	microns) microcapsules. Microcapsules under 10 microns in diameter tend to be readily
3	Microcapsules under 10 interests phagocytized and also tend to under go rapid spontaneous degradation due to phagocytized and also tend to under go rapid spontaneous degradation due to
4	phagocytized and also tend to under go op- their high surface to volume ratio. These smaller microcapsules would be well their high surface to volume ratio.
5	their high surface to volume rand. There is their pulse of antigen release could be suited for eliciting a primary response if their pulse of antigen release could be
6	suited for eliciting a primary response is
7	programmed to occur after phagocytosis. LITERATURE CITED
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PHASE III

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This phase of the invention relates to providing novel biocompatible and biodegradable microspheres for burst-free programmable sustained release of biologically active agents, inclusive of polypeptides, over a period of up to 100 days in an aqueous physiological environment. Potentially release period is capable of being further modulated beyond 100 days to about 365 days by careful selection of a blend of uncapped and end-capped biodegradable-biocompatible copolymer and molecular weights.

Several publications and patents are available for sustained
Soveral publications and patents are nolymers,
Several publications and paterns. 5 release of active agents from biodegradable polymers, 5 release of active agents from biodegradable polymers, 6 PLGA). Prior usages of
release of active/glycolides) (PLGA). Prior the use
release of active agents from biodegradable 1 release of active agents from biodegradable 1 particularly, poly(lactide/glycolides) (PLGA). Prior usages of particularly, poly(lactide/glycolides) (PLGA).
particularly, poly(lactide/glycolides) (PLGA) particularly, poly(lactide/glycolides) (PLGA) plga for controlled release of polypeptides have involved the use plga for controlled release of polypeptides have involved the use
of molar ratios of lactide/9-1
legular weights <20,000. In agreeus layer to
molecular ways
utilized fillers or additives in the interest and/or to improve the stability and encapsulation efficiency and/or to
improve the stability and encapsulation error improve the stability and encapsulation error improve the stability and encapsulation error in the advanced layer, thereby modulating increase the viscosity of the aqueous layer, thereby modulating increase the viscosity of the biologically active agent or
increase the Viscosia the biologically active agent of
increase the viscosity of the aqueous 12 increase the viscosity of the aqueous 13 polymer hydrolysis and the biologically active agent or 13
polypeptide release.
E DIGA Copolymers were end-
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exhibited a low to men in the first 24 hours after placement in the
exhibited a low to moderate burst release or exhibited a low to moderate burst release or entrapped polypeptide in the first 24 hours after placement in an entrapped polypeptide in the first 24 hours after placement in an
agueous physiological environment agueous phase.
aqueous physiological environment. It is aqueous phase. 22 are due to the use of fillers in the inner aqueous phase. 23 Further, a 1-month release of polypeptide is known with the use
LL SALPASE SE
23 Further, a 1-month relation of Mw <20,000. 24 of a 75/25 co-polymer of PLGA of Mw <20,000.
of a 75/25 co-polymer of PLGA of Mw 120,000 24 of a 75/25 co-polymer of PLGA of Mw 120,000 Investigations in controlled release research has been 100 to 2 month delivery system 100 to
Investigations in controlled release Investigations in controlled release The second in the second
26 proceeding especially to the

for biologically active agents or polypeptides using poly(lactide/glycolide) polymers. However, most of these systems have one or more of the following problems: Poor encapsulation efficency and large 'burst release' followed by an intermediate 'no release' or 'lag phase' until the polymer degrades. In general, release from these polymers occur over a period from about 4 weeks to about several months. In addition, in order to achieve this release a 50/50 copolymer of MW > 30,000 or a 75/25 copolymer of Mw > 10,000 are employed which often results in residual polymer remaining at the site of administration long after the release of active core.)

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This invention provides biocompatible and biodegradable microspheres that have been designed for novel, burst free, 3 programmable sustained release of biologically active agents, including polypeptides over a period of up to 100 days in an 5 6 aqueous physiological environment. 7 Unlike currently available release systems, which rely on 8 the use of fillers/additives such as gelatin, albumin, dextran, pectin, polyvinyl pyrrolidone, polyethylene glycol, sugars, etc., 0 $_{
m 12}$ and are still prone to low encapsulation efficiencies and "burst 13 effects", this invention achieves high encapsulation and "burst- $_{
m 14}$ free" release without the use of any additive. In this invention, burst-free, programmable sustained release is achieved through 16 the use of a unique blend of the 'uncapped' and end-capped forms 17 of poly(lactide/glycolide) polymer in the molecular weight range $_{18}$ of 2,000 to 60,000 daltons. In general, microspheres described in this invention are 19 21 produced by a unique emulsification technique wherein an imner) 22 water-in-oil (w/o) emulsion is stabilized by dispersing ima 23 solvent-saturated aqueous phase containing an emulsion 24 stabilizer. A ternary w/o/w emulsion is then formed by emulsifying the above w/o emulsions in an external pre-coded 26 aqueous phase containing an o/w emulsifier. Essentially, the

inner w/o emulsion is comprised of an aqueous layer containing from - 2 to about 20% (w/w) of the active agent to be entrapped 1 and an oil layer containing poly(lactide/glycolide) copolymer in 2 concentrations ranging from - 5 to about-- 50% (w/w oil phase). 3 The copolymer includes molecular weight ranging from 2,000 to 4 about 60,000 daltons, with molar composition of lactide/glycolide 5 from 90/10 to 40/60 and a blend of its uncapped and end-capped 6 forms in a ratio of 100/0 to 1/99. Very high encapsulation 7 efficiencies of about 80 to 100% are achieved depending on 8 9 polymer molecular weight and structural form. Programmable release of active core over variable durations 0 between 1-100 days is achieved by a judicious selection of 11 process parameters such as polymer concentration, peptide 12 concentration and the aqueous/oil phase ratio. 13 This invention is particularly suitable for high 14 encapsulation efficiencies and burst-free, continuous 5 programmable release of polypeptides of molecular weights ranging 16 from 1,000 to about 250,000 daltons, and also other biologically 17 active agents over a period of 1-100 days. A uniqueness of the 18 invention is that when using a 100/0 blend of the uncapped and 19 capped polymer, the final phase of active core release is) concurrent with the complete solubilization of the polymer to 21 innocuous components, such as lactic and glycolic acids. This is 22 a significant advantage over the currently available 30 day -23 release systems wherein a major regulatory concern is about 24 toxicity of residual polymer at the site of administration, long 26

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This invention relates to the design of biocompatible and 7 biodegradable microspheres for novel, programmable sustained 8 release of biologically active agents, including polypeptides 9 over a period of up to 100 days in an aqueous physiological ٥ environment with little or no burst release. 1 Unlike currently available release systems which rely on the 2 use of fillers/additives such as gelatin, albumin, dextran, 3 pectin, polyvinyl pyrrolidone, polyethylene glycol, sugars, etc., 4 and are still prone to low encapsulation efficiencies and "burst 5 effects", this invention achieves high encapsulation efficiency after release of the active core.

The microcapsules described in this invention are suitable 0 for administration via several routes such as parenteral 1 (intramuscular, subcutaneous), oral, topical, nasal, rectal and 3 vaginal routes.

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and 'burst-free' release without the use of any additive. In this invention, burst-free, programmable sustained release is achieved 1 through the use of a unique blend of the 'uncapped' and end-2 capped forms of poly (lactide/glycolide) polymer. .3 The 'uncapped' form refers to "poly(lactide/glycolide) with 4 free carboxyl end groups" which renders the polymer more 5 hydrophilic compared to the routinely used end-capped form. 6 Currently used 'end-capped' polymer hydrates between 4-12 weeks 7 depending on the molecular weight, resulting in an intermediate 8 'no release' or a 'lag phase'. The uncapped polymer hydrates 9 typically between 5 to 60 days depending on the molecular weight, 0 thus releasing its core continuously without a lag phase. A 11 careful blend of the two forms and appropriate molecular weights 12 and L/G ratios, results in a continuous release between 1 to 100 13 days. In addition, release within this time is programmable by a 14 judicious selection of process parameters such as polymer 5 concentration, peptide concentration and the aqueous/oil phase 16 17 The coploymer in this invention includes molecular weight ratio. 18 ranging from 2,000 to 60,000 daltons, a lactide/glycolide ratio 19 of 90/10 to 40/60 and a blend of the uncapped/capped forms in the) ratio of 100/0 to 1/99. The molecular weight of the polypeptide 21 may be in the range of 1000 to 250,000 daltons while that of ?2 other biologically active agents may range from 100 to 100,000 23 24 Microcapsules described in this invention are prepared by a daltons. j

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unique aqueous emulsification techinique which has been developed for use with the uncapped polymer to provide superior sphere 1 morphology, sphere integrity and narrow size distribution. This 2 is accomplished by first preparing an inner water-in-oil (w/o) by mixing the solutions of polymer in an organic solvent such as methylene chloride and the biologically active agent in water. 5 This is followed by stabilization of the w/o emulsion in a 6 solvent-saturated aqueous solution containing an o/w emulsifier 7 such as polyvinyl alcohol. A ternary emulsion is then formed by 8 9 emulsifying the w/o emulsion in an external aqueous phase containing the same emulsifier as above at concentrations ranging .0 from 0.25 - 1% w/v. Microcapsules are hardened upon solvent 11 removal by evaporation, rinsed to remove residual emulsifier and 12 lyophilized. Low temperature is used both at the time of primary 13 emulsification (w/o emulsion formation) and during the formation 14 of the final w/o/w emulsion to achieve stable emulsion and 5 16 superior sphere characteristics. In the context of the invention, a biologically active agent 17 is any water-soluble hormone drugs, antibiotics, antitumor 18 agents, antiinflammatory agents, antipyretics, analgesics, 19 3 antitussives, expectorants, sedatives, muscle relaxants, antiepileptics, antiulcer agents, antidepressants, antiallergic 21 drugs, cardiotonics, antiarrhythmic drugs, vasodilators, 22 antihypertensives, diuretics, anticoagulants, antinarcotics, 23 24 and the agents listed in the summary of the invention section herein 5.

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	More precisely, applicants have discovered a
1	pharmaceutical composition and process with the following
2	pharmaceutical composition and i
3	itemized features:
4	1. A controlled release microcapsule pharmaceutical formulation, which
	may contain a pharmaceutically-acceptable adjuvant, for burst-free, sustained,
5	programmable release of a biologically active agent over a duration from 1-100 days,
6	biologically active agent of uncapped and end-capped
7	comprising an active agent and a blend of uncapped and end-capped
8	biodegradable poly(lactide/glycolide).
	and a paragoutical formulation of item 1, wherein the
9	2. The pharmaced trade of uncapped and biodegradable poly(lactide/glycolide) is a blend of uncapped and
10	blodegradable poly(1222 promise from 100/0 to 1/99.
11	capped forms, in ratios ranging from 100/0 to 1/99.
12	3. The microcapsules of items 1 or 2 wherein the copolymer
13	(lactide to glycolide L/G) ratio for uncapped and endcapped
14	polymer is 52/48 to 48/52.
.5	right research the copyright of items 1 or 2 wherein the copyright
	and end-capped polymer is 90/10 to 40/00
16	5 mbs microcapsules of items 1 or 2 or 3 of 4 microcapsules
17	molecular weight of the copolymer is between 2,000-60,000
18	molecular weight of the Copoly
19	daltons.
0	6. The microcapsules of items 1 or 2 or 3 or 4 or 5 wherein
21	the biologically active agent is a peptide or polypeptide.
	innegatives of item 6, wherein said pour
22	7. The microcapsulos histatin consisting of 12 amino acids and having a molecular
23	
25	weight of 1563.
5	weight of 1563. 8. The microcapsules of items 1 or 2 or 3 or 4 or 5 or 6
20	8. The microcapsules of the completely release histatin in characterized by the capacity to completely release histatin in

an aqueous physiological environment from 1-35 days with a 100/0 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 48/52 to 52/48, and a molecular weight <15,000.

- 9. The microcapsules of items 1 or 2 or 3 or 4 or 5 or 6 characterized by the capacity to completely release histatin in an aqueous physiological environment from 18-40 days with a 100/0 blend of uncapped and end-capped poly(lactide/glycolide) having a L/G ratio of 48/52 to 52/48 and a molecular weight range of 28,000-40,000.
- 10. The microcapsules of items 1 or 2 or 3 or 4 or 5 or 6 0
- characterized by the capacity to release up to 90% of the 1
- histatin in an aqueous physiological environment from 28-70 days
- with a 0/100 blend of uncapped and end-capped 2 3
- poly(lactide/glycolide) having a L/G ratio of 48/52 to 52/48 and
- a molecular weight range of 10,000-40,000 daltons. 4
- 11. The microcapsules of items 1 or 2 or 3 or 4 or 5 or 6 5
- characterized by the capacity to release up to 80% of histatin in 6
- an aqueous physiological environment from 56-100 days with a 7 8
- 0/100 blend of uncapped and end-capped poly(lactide/glycolide)
- having a L/G ratio of 75/25 and a molecular weight of \prec 15,000 9)
- .1

- 12. The microcapsules of items 7 or 8 or 9 or 10 or 11 daltons.
- having analogs of histatin with chain lengths of from 11-24 amino .2 3
- acids of molecular weights from 1,500-3,000 daltons and characterized by the following structures:
- 1. DSHAKRHHGYKRKFHEKHHSHRGY 6

	2. KRHHGYKRKFHEKHHSHRGYR
1	T T H G V K R K F H E K H H S H R
2	RKFHEKHHSHRGYR
3	4. 5. AKRHHGYKRKFH
4	5. AKRHHGYKRKFH
5	- '
6	7. KRHHGYKRKF
7 .	* D-amino acid
8	13. The microcapsules of items 1 or 2 or 3 or 4 or 5 wherein
9	the biologically active agent is a polypeptide Leutinizing
10	hormone releasing hormone (LHRH) that is a decapeptide of
11	molecular weight 1182 in its acetate form, and having the
12	structure:
13	p- E H W S Y G L R P G
14	14. The microcapsule of items 6 or 7 or 8 or 9 or 10 or 11
5	or 12 or 13 having a molecular weight of from 1,000 to 250,000
16	or 12 or 13 having a molecular weight
17	daltons. 15. The microcapsules of items 6 or 7 or 8 or 9 or 10 or 11
18	15. The microcapsules of Items of variable rates and
19	or 12 or 13 or 14 wherein release profiles of variable rates and
၁	durations are achieved by blending uncapped and capped
21	microspheres as a cocktail in variable amounts.
22	microspheres as a cocktair and microspheres and microspheres and microspheres and microspheres as a cocktair and microspheres and microspheres and microspheres and microspheres and microspheres are a cocktair and microspheres and microspheres are a cocktair and microspheres and microspheres and microspheres and microspheres are a cocktair and microspheres are a
23	or 12 or 13 or 14 wherein release of profiles of variable rates
24	or 12 or 13 or 14 wherein to a solution are achieved by blending uncapped and capped polymer and duration are achieved by blending uncapped and capped polymer
i	in different ratios within the same microshreres.
26	in different ratios within 17. The microcapsules of items 6 or 7 or 8 or 9 or 10 or 11

- or 12 or 13 or 14 or 15 or 16 wherein the entrapped polypeptide
- is any of the vaccine agents against enterotoxigenic E. coli 1 2
- (ETEC) such as CFA/I,CFA/II,CS1,CS3,CS6 and CS17 and other ETEC-3
- related enterotoxins. 4
 - 18. The microcapsules of items 6 or 7 or 8 or 9 or 10 or 11
- or 12 or 13 or 14 or 15 or 16 or 17 wherein the entrapped 5
- polypeptide consists of peptide antigens of molecular weight 6 7
- range of about 800-5000 daltons for immunization against 8
- enterotoxigenic E. coli (ETEC). 9
- 19. The microcapsules of items 1 or 2 or 3 or 4 or 5 wherein 10
 - said biologically active agents are selected from the group
- consisting of water-soluble hormone drugs, antibiotics, antitumor 11
- agents, anti inflammatory agents, antipyretics, analgesics, 12
- antitussives, expectorants, sedatives, muscle relaxants, 13
- antiepileptics, antiulcer agents, antidepressants, antiallergic 14 15
 - drugs, cardiotonics, antiarrhythmic drugs, vasodilators,
- antihypertensives, diuretics, anticoagulants, and antinamotics, 16
- in the molecular weight range of 100-100,000 daltons. 17
- 20. The microcapsules of items 1 or 2 or 3 or 4 or 5 or 6 or 18
- 7 or 8 wherein said biodegradable poly(lactide/glycolide) is in 19 0
- an oil phase, and is present in about 1-50% (w/w).
- 21. The microcapsules of items 1 or 2 or 3 or 4 or 5 er 6 or 21
- 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 wherein 22
- concentration of the active agent is in the range of 0.1 to about 23
- 60% (W/W). 5
- 22. The microcapsules of items 1 or 2 or 3 or 4 or 5 sr 6 or 26

8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 wherein a ratio of the inner aqueous to oil phases is about 1/4 to

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23. A process for preparing controlled release microcapsule 1/40(V/V). formulations characterized by burst-free, sustained, programmable release of biologically active agents comprising: Dissolving biodegradable poly (lactide/glycolide), in uncapped form in methylene chloride, and dissolving a biologically active agent or active core in water; adding the aqueous layer to the polymer solution and emulsifying to provide an inner water-in-oil (w/o) emulsion; stabilizing the w/o emulsion in a solvent-saturated .0 aqueous phase containing a oil-in-water (o/w) emulsifier; adding . 1 said w/o emulsion to an external aqueous layer containing oil-in-. 2 water emulsifier to form a ternary emulsion; and stirring the .3 resulting water-in-oil-in-water (w/o/w) emulsion for sufficient .4 time to remove said solvent, and rinsing hardened microcapsules 5 with water and lyophilizing said hardened microcapsules. .6 24. A process for preparing controlled release microcapsule .7 formulations characterized by burst-free, sustained, programzble .8 .9 release of biologically active agents comprising: dissolving biodegradable poly(lactide/glycolide) in end-0 capped form in methylene chloride, and dissolving a biologically !1 active agent or active core in water; adding the aqueous layer to 2. the polymer solution and emulsifying to provide an inner water-:3 in-oil emulsion; stabilizing the w/o emulsion in a solvent-!4 saturated aqueous phase containing a oil-in-water (o/w) 5

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emulsifier; adding said w/o emulsion to an external aqueous layer

containing oil-in-water emulsifier to form a ternary emulsion; ı

and stirring a resulting water-in-oil-water (w/o/w) emulsion for 2

sufficient time to remove said solvent; and rinsing hardened

microcapsules with water; and lyophilizing said hardened 5

microcapsules.

25. The process of items 23 or 24 wherein a solvent-6 7

saturated external aqueous phase is added to emulsify the inner 8

w/o emulsion prior to addition of the external aqueous layer, to

provide microcapsules of narrow size distribution range between

0.05-500µm. 11

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26. The process of items 23 or 24, wherein a low temperature

of about 0-4°C is provided during preparation of the inner w/o 12

emulsion, and a low temperature of about 4-20°C is provided 13

during preparation of the w/o/w emulsion to provide a stable 14

emulsion and high encapsulation efficiency. 16

27. The process of items wherein a 100/0 blend of uncapped

and end-capped polymer is used to provide release of the active 17

core in a continous and sustained manner without a lag phase. 18

28. The microcapsules of items 6, wherein, when the

entrapped polypeptide is active at a low pH, such as LHRH,

adrenocorticotropic hormone, epidermal growth factor, calcitomin 21

released polypeptide is bioactive. 23

29. The microcapsules of items 6 or 7 or 8 or 9 or 10 or 11, 24

wherein, when entrapped peptide such as histatin is inactive at a

26 low pH, a pH-stabilizing agent of inorganic salts are added to

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the inner aqueous phase to maintain biological activity of the ١ released peptide.

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- 30. The microcapsules of items 6 or 7 or 8 or 9 or 10 or 11 wherein, when entrapped polypeptide such as histatin is inactive at a low pH, a non-ionic surfactant such as polyoxyethylene sorbitan fatty acid esters (Tween 80, Tween 60 and Tween 20) and polyoxyethylene - polyoxypropylene block copolymers (Pluronics) is added to the inner aqueous phase to maintain biological 7 3 activity of the released polypeptide.
- 31. The microcapsules of items 29, wherein placebo spheres 7 loaded with the pH-stabilizing agents are coadministered with 2 polypeptide-loaded spheres to maintain the solution pH around the 11 microcapsules and preserve the biological activity of the 12 released peptide in instances where the addition of pH-13 stabilizing agents in the inner aqueous phase is undesirable for 14 the successful encapsulation of the acid pH sensitive 5 16 polypeptide. 17
 - 32. The microcapsules of item 30 wherein placebo spheres loaded with non-ionic surfactant are coadministered with 18 polypeptide-loaded spheres to maintain biological activity of the 19 released peptide where the addition of non-ionic surfactants in 1 the inner aqueous phase is undesirable for successful 21 22 encapsulation of the acid pH sensitive polypeptide. 23
 - 33. The microcapsules of items 1 or 2 or 3 or 4 or 5 or 6 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 comprising 24 a blend of uncapped and capped polymer, wherein complete 26

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solubilization of the copolymer leaves no residual polymer at the L site of administration and occurs concurrently with the complete 2 release of the entrapped agent. 3 34. A process of using microcapsules of items 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or 5 16 or 17 or 18 or 19 or 20 for human administration via parenteral routes, such as intramuscular and subcutaneous. 35. A process of using microcapsules of items 1 or 2 or 3 or 3 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 or 18 or 19 or 20 for human administration via topical 0 route. 11 36. A process of using microcapsules of items 1 or 2 or 3 or ι2 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or ι3 16 or 17 or 18 or 19 or 20 for human administration via oral ί4 5 routes. 37. A process of using microcapsules of items 1 or 2 or 3 or .6 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or .7 16 or 17 or 18 or 19 or 20 for human admininstration via nasal, .8 transdermal, rectal, and vaginal routes. .9 !1 :2

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Conservation of bioactivity of polypeptides As the polymer degrades rapidly, there is a precipitous 2 drop in pH accompanied by the release of soluble oligomers in the 3 microenvironment which may affect the biological activity of acid 4 pH-sensitive peptides/proteins. In such instances, biological 5 activity can be maintained by the use of inorganic salts or 6 buffering agents in the inner aqueous phase codissolved with the 7 8 The following unique advantages are characteristics of this peptide. 9 0. 1. Burst-free, prolonged, sustained release of polypeptides invention: 11 and other biologically-active agents from biocompatible and biodegradable 12 microcapsules up to 100 days in an aqueous physiological environment without 13 14 the use of additives in the core. 2. Release of active core programmable for variable 5 durations over 1-100 days, by using a blend of uncapped and 16 capped polymer of different molecular weights and copolymer 17 18 ratio, and by manipulating the process parameters. 3. Complete release of the active core is concurrent with 19 complete solubilization of the carrier polymer to innocuous) 21

21 complete release of the active core is concurrent with
22 complete solubilization of the carrier polymer to innocuous
23 components, such as lactic and glycolic acids, especially when
24 using a 100/0 blend of uncapped/capped polymer. This is of
25 tremendous significance, as most biodegradable polymers currently
26 used for 1-30 day delivery, do not degrade completely at the end
27 of the intended release duration, thereby causing serious concern

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1 c. regulatory authorities on the effects of residual polymer at the site of administration.

4. Ease of administration of the microcapsules in various dosage forms via several routes, such as parenteral

5 (intramuscular and sucutaneous), oral, topical, nasal, vaginal, etc.

The hydrophilic homo-and co-polymers based on D,L-lactide
and glycolide contains hydrophilic adjusted homo-and co-polymers
with free carboxylic end groups, and is characterized by the
formula:

16
17 $(C_3H_4O_2)_n(C_2H_2O_2)_m$ n:m = 1:1

Wherein Z= Molecular Weight/130; for example Z=92 for Mw 19 12,000 and 262 for Mw 34,000.

While the molar ratio of the lactide to glycolide may vary,
it is most preferred that the lactide to glycolide copolymer
ratio be 50:50.

Reference is now made to FIG. 48 which depicts a blood-drug

24 concentration versus time graph that shows conventional drug

25 administration using a series of dosages compared to an ideal

26 controlled release system. Unfortunately, many drugs have a

blending of the two forms in a single formulation comprising different ratios of uncapped to capped polymer, would significantly influence the polymer hydration and hence release of the active core thereby providing release curves of any desirable pattern. Manipulation of polymer hydration and degradation resulting in modulation of release of active core is achieved by the addition of uncapped polymer to end-capped polymer in amounts as low as 1% up to 100%. .0

While referring to Table 14 in conjunction with FIG.50 , it can be seen that the cumulative Histatin release from PLGA 1 microspheres from several batches prepared using 50/50 and 75/25 2 uncapped and end-capped, polymer modulates release between 1 to 3 100 days by varying the process parameters. 1-35 days by uncapped 50/50, 18-56 days by capped 50/50 and 56-100 days by capped 5 5 In referring to FIG.51, a view is provided through a 75/25. 7 scanning electron micrograph of PLGA microspheres designed for a 3 one to two month release system prepared using end-capped polymer) 0 of Mw 30-40k daltons. 11 FIG. 52 depicts the cumulative Histatin release from PLGA 12 microspheres, in which the release profiles are from several 13 batches prepared using 50/50, uncapped and capped polymer, and 14 varying the process parameters to modulate release between 28 to 5 16 Figure 53 represents cumulative Histatin release from PLGA 60 days. 17 microspheres --- these combined release profiles are from several 18 batches prepared using 50/50 uncapped and capped polymer, and 19 varying the process parameters to modulate release between 1-60 0 21 In the context of the invention, a biologically active agent days. ?2 is any water-soluble antibiotics, antitumor agents, antipyretics 13 analgesics, anti-inflammatory agents, antitussives, expectorants, 14 sedatives, muscle relaxants, anti epileptics, antiulcer agents, 5 16

	cardiotonics,
1	anti-depressants, anti-allergic drugs, cardiotonics,
2	anti-depressants, anti-depress
3	antiarrhythmics drugs, vasodifatory drugs, anti-narcotics, etc. diuretics, anticoagulants, hormone drugs, anti-narcotics, etc.
4	enell eneralited re-
5	In general, "burst free sussession of sold of
6	in the second of
7	in the context of this invention does not be context.
8	
9	In general, the approaches for designing the biologically
10	In general, the approximation active agents encapsulated in the uncapped and combination
11	uncapped/end-capped PLGA microspheres and characteristics of
12	uncapped/end-capped Flor missis these encapsulants are briefly set forth below as follows:
13	1. Providing PLGA microspheres of surface morphologies using 50/50 uncapped and capped polymers of Mw - 8-40K daltons as shown
14 15	in Flas. 40 and li
16 17 18	malage of a polypeptide, and uncapped
19	and capped polymer
50	50/50 and 73/23
21 22	50/50 and 75/25. For example, design of a 1-12 week bioactive compound release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using PLGA with the following release system is achieved using places and the following release system is achieved using places are achieved using places and the following release system is achieved using places and the following release system is achieved using places.
23 24	
! 5	
26 27	1. Polymer molecular weight: -about 2-60K daltons -atio (L/G):
27 28	a complymer molar facto (2/5/
29	_ 90/10 to 40/0-
0	3. Polymer end groups: - uncapped and /or end-capped - uncapped and /or end-capped
3)	and combining judiciously within the following parameters:
33	and combining justing
3 [,] 5	4. Polymer concentration - from 5 to 50% - phase ratio:
_	6
3	_ 1:5 to **=* * ' '
3	. PAM Z CO GATE
U	·

and by using the unique aqueous emulsification method described in the invention.

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The uniqueness and novelty of invention may generally be summarized in a brief way as follows:

- 1. Use of uncapped poly(lactide/glycolide) to achieve burstfree, continuous, sustained, programmable release of biologically
- 2. Use of a unique aqueous emulsification system to achieve active agents over 1-100 days. superior microsphere characteristics such as uniform sphere morphology and narrow size distribution.
- 3. Burst-free, prolonged, sustained release of polypeptides and other biologically actice agents from biocompatible and biodegradable microcapsules up to 100 days in an aqueous physiological environment without the use of additives in the
- 4. Release of active core programmable for variable durations over 1-100 days by using a blend of uncapped and capped polymer for different molecular weights and copolymer rations and
- 5. Complete release of the active core concurrent with manipulating the process parameters. complete solubilization of carrier polymer to innocuous components such as lactic and glycolic acids, especially when using a 100/0 blend of uncapped/capped polymer. This is of tremendous significance as most biodegradable polymers currently in use for 1-30 day delivery, do not degrade completely at the end of the intended release duration causing serious concern for regulatory authorities on the effects of residual polymer at the
- 6. Ease of administration of the microcapsules in various dosages forms via several routes such as parenteral (intramusclar site of administration. and subcutaneous), oral, topical, nasal, vaginal, etc.

The following examples are illustrative of, but not limitations upon the microcapsule compositions pertaining to this invention.

Example 12

Polylactide/glycolide (PLGA) microcapsules are prepared by a unique aqueous emulsification technique which has been developed for use with the uncapped polymer to provide superior sphere morphology, sphere integrity and narrow size distribution (See Figures 32 and 322). This is accomplished by dissolving the polymer in a chlorinated hydrocarbon solvent such as methylene chloride and dissolving the biologically active agent in water. A

w/o emulsion is then formed by mixing the solutions of polymer and the active agent by sonication, followed by emulsion stabilization in a solvent - saturated aqueous solution containing polyvinyl alcohol. A ternary emulsion is then formed by emulsifying the w/o emulsion in an external, pre-cooled aqueous phase containing polyvinyl alcohol (0.25 - 1% w/v). Microcapsules are hardened upon removal of solvent by evaporation, rinsed to remove any residual emulsifier, and then lyophilized.

Table14 lists the microcapsule compositions, Nos. 1-21 thus prepared, consisting of a biologically active polypeptide, Histatin (composed of 12 amino acids and a molecular weight of 1563) and blends of uncapped and capped polymer of ratios 100/0 to 1/99, and having a lactide/glycolide ratio of 90/10 to 40/60, and a molecular weight range between 2000 to 60,000 daltons.

Example 13

Microcapsule compositions are prepared as described in Example 1 wherein the copolymer L/G ratio is 48/52 to 52/4E, and the ratio of uncapped/capped polymer is 100/0. The active core is Histatin (Mw 1563), the polymer molecular weight is < 15,000 and the polymer concentrations vary from 7% to - 40% w/w. Compositions 1,2,4 12-14 and 16-18 are listed in Table14.

Release profiles of the active core from the compositions in an aqueous physiological environment, such as phosphate-buffered saline, pH 7.0 maintained at 37 ± 1°C are plotted as cumulative percentage release versus time, and presented in Figure 50

Burst-free, variable release from 1-35 days is achieved by varying the polymer concentration from 7 to - 40% w/w in the oil phase.

Example 14

Microcapsule compositions are prepared as described in Example 2, wherein the aqueous /oil ratio is varied from 1/4 to 1/20 (v/v). Compositions 1,2,4 and 12 are listed in Table 14

Release profiles of the active core from the compositions in an aqueous physiological environment described in Example are plotted as cumulative percentage release versus time, and presented in Figure 50.

Burst-free, continuous release from 1-35 days, with different onset and completion times are achieved by selecting different w/o ratios in the inner core.

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Example 15

Microcapsule compositions are prepared as described in Example 2, wherein the polymer molecular weight is 28,000-40,000 and polymer concentrations vary from 5% to - 15% W/W. Compositions 19-21 are listed in Table 14.

Release profiles of the active core from the compositions in an aqueous physiological environment are described in Example 2 are plotted as cumulative percentage release versus time and presented in Figure 52.

Burst-free, variable release from 18-40 days is achieved by varying the polymer concentration.

Example 16

Microcapsule compositions are prepared as described in Example 2, wherein the ratio of uncapped/capped polymer is 1/99 and polymer concentrations vary between 5% to - 12% W/W. Compositions 10 and 11 are listed in Table14...

Release profiles of the active core from the compositions in an aqueous physiological environment are described in Example 2, and plotted as cumulative percentage release versus time and presented in Figure 50.

Burst-free, variable release from 28-70 days is achieved by varying the polymer concentration in the oil phase.

Example 17

Microcapsule compositions are prepared as described in Example 5, wherein polymer molecular weight is 28,000-40,000 and polymer concentrations vary between 5% to - 12% w/w. Compositions

5 and 6 are listed in Table 14. Release profiles of the active core from the compositions in an aqueous physiological environment are described in Example 2 and are plotted as cumulative percentage release versus time, and presented in Figure 52.

Burst-free, variable release from 28-70 days is achieved by varying the polymer concentration.

Example 18

Microcapsule compositions are prepared as described in Example 6, wherein the aqueous/oil ratio varies between 1/5 to 1/25 (V/V). Compositions 3 and 7 are listed in Table 14.

Release profiles of the active core from the compositions in an aqueous physiological environment are described in Example 2, and plotted as cumulative percentage release versus time, and

Burst-free, variable release from 28-70 days is achieved by presented in Figure 52 varying the aqueous/oil ratios.

Example 19

Microcapsule compositions are prepared as described in Example 5, wherein the copolymer ratio is 75/25 and polymer concentrations vary between 5% to - 25% W/W. Compositions 8 and 9

Release profiles of the active core from the compositions in are listed in Table 1. an aqueous physiological environment are described in Example 2, and are plotted as cumulative percentage release versus time, and

Burst-free, variable release from 56->50 days is achieved by presented in Figure 50. varying the polymer concentration in the oil phase.

Example 20

Microcapsule compositions are described in Example 2, wherein the active core is leutinizing hormone releasing hormone (LHRH, a decapeptide of molecular weight 1182) and the polymer concentration is -40% W/W. Release profiles of the active core from the composition in an aqueous physiological environment is described in Example 2, and is plotted as cumulative percentage release versus time, and presented in Figure 54

Burst-free, continuous and complete release is achieved within 35 days, similar to Histatin acetate.

Example 21

Microcapsule compositions are prepared as described in Example 2, wherein an additive such as sodium salt (carbonate or bicarbonate) is added to the inner aqueous phase at concentrations of 1-10% W/W to maintain the biological activity of the released polypeptide.

Burst-free, variable release from 1-28 days is achieved similar to Examples 2 & 3, and the released polypeptide is biologically active until 30 days, due to the presence of the sodium salt.

Example 22

Microcapsule compositions are prepared as described in Example 2, wherein an additive such as a nonionic surfactant, polyoxyethylene/polyoxypropylene block copolymer (Pluronics F68 and F127) is added to either the inner oil or the aqueous phase at concentrations from 10-100% w/w, to maintain the biological activity of the released polypeptide.

Burst-free, continuous release from 1-35 days is achieved similar to Examples 2 & 3, and the released polypeptide is bioactive due to the process of the surfactant bioactive due to the presence of the surfactant.

Example 23

Cumulative histatin release from the microcapsule compositions described in Examples 1 through 11 and release profiles plotted in Figures49 and 50 show the burst-free, programmable peptide release for variable duration from 1-100 days. Virtually any pattern of cumulative release is achievable over a 100 day duration by a judicious blending of several compositions, as shown in Figure 53.

- What we claim is:
- 1. A composition for the burst-free, sustained, programmable 1
- release of active material(s) over a period from 1-100 days, 2
- which comprises: (1) An active material and (2) A carrier which 3
- may contain pharmaceutically-acceptable adjuvant, comprised of a
- blend of uncapped and end-capped biodegradable-biocompatible 5 6
- 7
- 2. The composition of Claim 1 wherein the polymeric substance is copolymer. . 8
 - poly(lactide/glycolide).
 - 3. The composition of Claim 2, wherein the 10
 - poly(lactide/glycolide) is a blend of uncapped and end-capped 11
 - forms, in ratios ranging from 100/0 to 1/99. 12
 - The composition of Claim 3 wherein the copolymer (lactide to
 - glycolide L/G) ratio for uncapped and end-capped polymer is 90/10 13
 - 15
 - 5. The composition of Claim 4 wherein the copolymer (lactide to to 40/60. 16
 - glycolide L/G) ratio for uncapped and end-capped polymer is 48/52 17
 - 18
 - 6. The composition of Claim 2 wherein the molecular weight of to 52/48. 19
 - the copolymer is between 2,000-60,000 daltons.
 - The composition of Claim 3 wherein the active material is 21
 - biologically active agent. 22
 - The composition of Claim 7 wherein the agent is selected from
 - the group consisting essentially of antibacterial agents; 23 24
 - peptides; polypeptides; antibacterial peptides; antimycobacterial
 - agents; antimycotic agents; antiviral agents; antiparastic 25

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agents; antifungal; hormonal peptides; cardiovascular agents; hormonal peptides; cardiovascular agents; narcotic antagonists; 1 analgesics; anesthetics; insulins; steroids including HIV 2 therapeutic drugs (including protease inhibitors) and AZT; 3 estrogens; progestins; gastrointestinal therapeutic agents; non-4 steroidal anti-inflammatory agents; parasympathoimetic agents; 5 psychotherapeutic agents; tranquilizers; decongestants; sedative-6 hypnotics; non-estrogenic and non-progestional steroids; 7 sympathomimetic agents; vaccines; vitamins; nutrients; anti-8 migraine drugs; electrolyte replacements; ergot alkaloids; anti-9 inflammary agents; prostaglandins; cytotoxic drugs; antigens; 10. antibodies; enzymes; growth factors; immunomodulators; 11 pheromones; prodrugs; psychotropic drugs; nicotine; antiblood 12 clotting drugs; appetite suppressants/stimulants and combinations 13 thereof; contraceptive agents include estrogens such as diethyl 14 silbestrol; 17-beta-estradiol; estrone; ethinyl estradiol; 15 mestranol; progestins such as norethindrone; norgestryl; 16 ethynodiol diacetate; lynestrenol; medroxyprogesterone acetate; 17 dimethisterone; megestrol acetate; chlormadinone acetate; 18 norgestimate; norethisterone; ethisterone; melentate; 19 norgestimate; norethisterone; ethisterone; melengestrol; 20 norethynodrel; and spermicidal compounds such as 21 nonyphenoxypolyoxyethylene glycol; benzethonium chloride; 22 chlorindanol; include gastrointestinal therapeutic agents such as 23 aluminum hydroxide; calcium carbonate; magnesium carbonate; 24 sodium carbonate and the like; non-steroidal antifertility 25 26

agents; parasympathomimetic agents; psychotherapeutic agents; major tranquilizers such as chloropromaquine HCL; clozapine; 1 mesoridazine; metiapine; reserpine; thioridazine; minor 2 tranquilizers such as chlordiazepoxide; diazepam; meprobamate; 3 temazepam and the like; rhinological decongestants; sedative-4 hypnotics such as codeine; phenobarbital; sodium pentobarbital; 5 sodium secobarbital; other steroids such as testosterone and 6 testosterone propionate; sulfonmides; sympathomimetic agents; 7 vaccines; vitamins and nutrient such as the essential amino 8 acids; essential fats; anti-HIV agents; including AZT; 9 antimalarials such as 4-aminoquinolines; 8 aminoquinolines; 10 pyrimethamine; anti-migraine agents such as mazindol; 11 phentermine; anti-Parkinson agents such as L-dopa; antispasmodics 12 such as atropine; methscopolamine bromide; antispasmodics and 13 anticholingeric agents such as bile therapy; digestants; enzymes 14 and the like; antitussives such as dextromethorphan and 15 noscapine; bronchodilators; cardiovascular agents such as anti-16 hypertensive compounds; Rauwolfia alkaloids; coronary 17 18 vasodilators; nitroglycerin; organic nitrites; pentaerythriotetranitrate; electrolyte replacements such as 19 potassium chloride; ergotalkaloids such as ergotamine with and 20 21 without caffein; hydrogenated ergot alkaloids; dihydroergocristina methanesulfate; dihydroergocornine 22 methanesulfonate; dihydroergokroyptine methaneusulfate and 23 combinations thereof; alkaloids such as atropine sulfate; 24 Belladonna; hyoscine hydrobromide; analgesics; narcotics such as 25

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codeine; dihydrocodienone; meperidine; morphine; non-narcotics such as salicylates; aspirin; acetaminophen; and d-propoxyphene; 1 antibiotics such as the cephalosporins including ceflacor and 2 cefuroxime; chloranphenical; gentamicin; Kanamycin A. Kanamycin B; the penicillins; ampicillin; amoxicillin; streptomycin A; 4 antimycin A; chloropamtheniol; metromidazole; oxytetracycline 5 penicillin G; the tetracyclines; including minocycline; fluoro-6 quinolones including ciprofloxacin; ofoxacin; macrolides 7 including clarithromycin; frythromycin; aminioglycosides 8 including gentamicin; amikacin; tobramycin and kanamycin; beta-9 lactams including ampacillin; polymyxin-B; amphotercin-B; 10 11 aztrofonam; chloramphenicol; fusidans; lincosamides; metronidazole; nitro-furantion; imipenem/cilastin; quinolones; 12 systemic antibodies including rifampin; polygenes; sulfunamides; 13 trimethoprim; glycopeptides including vancomycin; teicoplanin and 14 imidazoles; anti-cancer agents; including anti-kaposi's sarcoma; 15 16 anti-convulsants such as mephenytoin; phenobarbital; 17 trimethadione; anti-emetics such as triethylperazine; 18 antihistamines such as chlorophinazine; dimenhydrinate; diphenhydramine; perphenazine; tripelennamine and the like; anti-19 inflammatory agents such as hormonal agents; hydrocortisons; 20 prednisolone; prednisone; non-hormonal agents; allopurinol; for 21 claims water-soluble hormone drugs; antibiotics; antitumor agents; anti inflammatory agents; antipyretics; analgesics; 23 24 antitussives; expectorants; sedatives; muscle relaxants; antiepileptics; anticulcer agents; antidepressants; antiallergic 25 26

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drugs; cardiotonics; antiarrhythmic drugs; vasodilators; antihypertensives; diuretics; anticoagulants; and antinarcotics; 1 in the molecular wight range of 100-100;000 daltons; 2 indomethacin; phenylbutazone; prostaglandins; cytotoxic drugs 3 such as thiotepa; chloramucil; cyclophosphamide; melphala; 4 nitrogen mustard; methotrexate; antigens such as proteins; 5 glycoproteins; synthetic peptides; carbohydrates; synthetic 6 polysaccharides; lipids; glycolipids; lipopolysaccharides(LPS); 7 synthetic lipopolysaccharides and with or without attached 8 adjuvants such as synthetic muramyl dipeptide derivatives; 9 antigens of such microorganisms as Neisseria gonorrhea; 10 Mycobacterium tuberculosis; Picarinii Pnfumonia; Herpes virus 11 (humonis types 1 and 2); Herpes zoster; Candidia albicans; 12 Candida tropicalis; Trichomonas vaginalis; Haemophilus vaginalis; 13 Group B streptoccoccus ecoli; Microplasma hominis; Hemophilus 14 ducreyi; Granuloma inguimale; Lymphopathia venerum; Treponema 15 palidum; Brucela aborus Brucela meitensis Brucela suis; Brucella 16 canis Campylobacter fetus; Campylobacer fetus intesinalis; 17 Leptospira pomona; Listeria monocytogenes; Brucella ovis; Equine 18 herpes virus 1; Equine arteritis virus; IBR-IBP virus; Chlamydia 19 psittaci; Trichomonas foetus; Taxoplasma gondii; Escherichia 20 coli; Actinobacillus equili; Salmonella abortus ovis. Salmonella 21 abortus eui; Pseudomonas aeruginosa; Corynebacterium equi; 22 Corynebacterium pyogenes; Actinobaccilus seminis; Mycoplassa 23 bovigenitalium; Aspergilus fumigatus; Absidia ramosa; Trypanosoma equiperdum; Babesia cabali; Clostridium tetani; 26

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antibodies which counteract the above microorganisms; and enzymes including ribonuclease; neuramidinase; trypsin; glycogen 1 phosphorylase; sperm lactic dehydrogenase; sperm hyaluronidase; 2 adenossinetriphosphase; alkaline phosphatase; alkaline phospha 3 esterase; amino peptides; typsin chymotrypsin amylase; 4 muramidase; acrosomal proteinase; diesterase; glutamic acid 5 dehydrogense; succunic and dehydrogenase; beta-glycophosphatase 6 lipase; ATP-ase alpha-peptate gamma-glutamyiotranspeptidase; 7 sterold-beta-ol-dehydrogenase; DPN-di-aprorase; and combinations 8 9 The composition of Claim 8 wherein the agent is selected from thereof. 10 the group consisting essentially of antibacterial agents; 11 antibacterial peptides; antimycobacterial agents; antimycotic 12 agents; antiviral agents; antiparasitic agents; antifungal; 13 hormonal peptides; cardiovascular agents; narcotic antagonist; 14 analgesics; anesthetics; vaccines; insulins; HIV therapeutic 15 drugs (protease inhibitors); estrogens; progestins; 16 gastrointestinal therapeutic agents; non-steroidal anti-17 inflammatory agents; parasympathoimetic agents; psychotherapeutic 18 agents; tranquilizers; decongestants; sedative-hypnotics; non-19 estrogenic and non-progestional steroids; sympathomimetic agents; 20 vaccines; vitamins; nutrients; anti-malarial compounds; asti-21 migraine drugs; electrolyte replacements; ergot alkaloids; 22 analgetics; non-narcotics; anti-cancer agents; anticonvulsmts; 23 anti-emetics; antihistamines; anti-inflammary agents; 24 prostaglandins; cytotoxic drugs; antigens; antibodies; enques; 25 26

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- growth factors; immunomodulators; pheromones; prodrugs;
- psychotropic drugs; appetite suppresants/stimulants; and 1
- combinations thereof. 3
- 10. The composition of Claim 8 wherein the agent is a peptide or 4
- polypeptide. 5
- The composition of Claim 10 wherein the agent is a poly 6
- 7
- The composition of Claim 11 wherein the molecular weight of peptide. 8
- the polypeptide is between 1,000-250,000 daltons.
- The composition of Claim 12 wherein the polypeptide is
- histatin consisting of 12 amino acids and having a molecular 10 11
- weight of 1563.
- 14. The composition of Claim 1 characterized by the capacity to
- completely release histatin in an aqueous physiological 13 14
- environment within from 1 to 40 days with a 100/0 blend of
- 16 uncapped and end-capped poly(lactide/glycolide) having a L/G 15
- ratio of 48/52 to 52/48, and a molecular weight less than 15,000.
- 15. The composition of Claim 14 wherein the histatin can be 17
- completely released within 18 to 40 days and the molecular weight 18
- of the poly(lactide/glycolide) is within the range of 28,600 to
- 16. The composition of Claim 2 characterized by the capacity to
- release up to 90% of the histatin in an aqueous physiological
- environment from 28-70 days with a 1/99 blend of uncapped and
- end-capped poly(lactide/glycolide) having a L/G ratio of 48/52 to 25
- 52/48 and a molecular weight range of 10,000-40,000 daltoms.

- 17. The composition of Claim 2 characterized by the capacity to 1
- release up to 80% of histatin in an aqueous physiological 2
- environment from 56-100 days with a 1/99 blend of uncapped and 3
- end-capped poly(lactide/glycolide) having a L/G ratio of 75/25 4
- and a molecular weight of less than 15,000 daltons. 5
- 18. The composition of Claim 13 having analogs of histatin with
- chain lengths of from 11-24 amino acids of molecular weights from 6 7
- 1,500-3,000 daltons and characterized by the following 8
- structures: 9
- 1. DSHAKRHHGYKRKFHEKHHSHRGY
- KRHHGYKRKFHEKHHSHRGYR 10 2.
- KRHHGYKRKFHEKHHSR 11 3. 12
- RKFHEKHHSHRGYR 4.
- 13 AKRHHGYKRKFH 5.
- 14 *A K R H H G Y K R K F H 6. 15
- KRHHGYKRKF 7. 16
- *D-amino acid 17
- 19. The composition of Claim 10 wherein the biologically active
- agent is a polypeptide Leutinizing hormone releasing hormone 18 19
- (LHRH) that is a decapeptide of molecular weight 1182 in its
- acetate form, and having the structure: 20 21
- P-EHWSYGLRPG 22
- The composition of Claim 13 having a molecular weight of 23
- from 1,000 to 250,000 daltons. 24

- The composition of Claim 2 wherein release profiles of
- variable rates and durations are achieved by blending uncapped ı
- and capped microspheres as a cocktail in variable amounts. 2
- 22. The composition of Claim 2 wherein release of profiles of 3
- variable rates and duration are achieved by blending uncapped and
- capped polymer in different ratios within the same microspheres. 5 6
- The composition of Claim 12 wherein the entrapped
- polypeptide is any of the vaccine agents against enterotoxigenic 7 8
- E. coli (ETEC) selected from the group consisting of 9
- CFA/I,CFA/II,CS1,CS3,CS6 and CS17, ETEC-related enterotoxins, and 10
- combinations thereof. 11
- 24. The composition of Claim 23 wherein the entrapped 12
- polypeptide consists of peptide antigens of molecular weight 13
- range of about 800-5000 daltons for immunization against 14
- enterotoxigenic E. coli (ETEC). 15
- The composition of Claim 24 wherein the entrapped 16
- polypeptide is selected from the group consisting essentially of 17
- an antigenic synthetic peptide containing CFA/I pilus protein T-18
- cell epitopes; B-cell epitopes, or mixtures thereof. 19
- 26. The composition of Claim 24 wherein the 20
- poly(lactide/glycolide) is a blend of uncapped and end-capped 21
- forms, in ratios ranging from 48/52 to 52/48. 22
- 27. The composition of Claim 7 wherein said agent are selected 23
- from the group consisting of water-soluble hormone drugs, 24
- antibiotics, antitumor agents, anti inflammatory agents,
- antipyretics, analgesics antitussives, expectorants, sedatives, 26

- muscle relaxants, antiepileptics, antiulcer agents, 1
- antidepressents, antiallergic drugs, cardiotonics, antiarrhythmic 2
- drugs, vasodilators, antihypertensives, diuretics, 3
- anticoagulants, antinarcotics, in the molecular weight range of 4
- 100-100,000 daltons.
- 28. The composition of Claim 1 wherein said biodegradable
- poly(lactide/glycolide) is in an oil phase, and is present in 6 7
- about 1-50% (W/W). 8
- 29. The composition of Claim 28 wherein concentration of the 9
- active agent is in the range of 0.1 to about 60% (w/w). 10
- The composition of Claim 29 wherein a ratio of the inner 11
- aqueous to oil phases is about 1/4 to 1/40(v/v). 12
- The composition of Claim 11 wherein the entrapped 13
- polypeptide is active at a low pH, such as LHRH, 14
- adrenocorticotropic hormone, epidermal growth factor, calcitonin 15
- released polypeptide is bioactive. 16
- 32. The composition of Claim 11 when entrapped polypeptide such 17
- as histatin is inactive at a low pH, a pH-stabilizing agent of 18
- inorganic salts are added to the inner aqueous phase to maintain 19
- biological activity of the released peptide. 20
- 33. The composition of Claim 11 wherein when entrapped 21
- polypeptide such as histatin is inactive at a low pH, a non-ionic 22
- surfactant such as polyoxyethylene sorbitan fatty acid esters 23
- (Tween 80, Tween 60 and Tween 20) and polyoxyethylene -24
- polyoxypropylene block copolymers (Pluronics) is added to the 25

- inner aqueous phase to maintain biological activity of the 1
- released polypeptide.
- 34. The composition of Claim 32 wherein placebo spheres loaded
- with the pH-stabilizing agents are coadministered with
- polypeptide-loaded spheres to maintain the solution pH around the 5
- microcapsules and preserve the biological activity of the
- released peptide in instances where the addition of pH-stablizing 6 7
- agents in the inner aqueous phase is undesirable for the
- 9. successful encapsulation of the acid pH sensitive polypeptide.
- 10 35. The composition of Claim 33 wherein placebo spheres loaded
- ll with non-ionic surfactant are coadministered with polypeptide-
- 12 loaded spheres to maintain biological activity of the released
- 13 peptide where the addition of non-ionic surfactants in the inner
- 14 aqueous phase is undesirable for successful encapsulation of the
- 15 acid pH sensitive polypeptide.
- 16 36. The composition of Claim 1 comprising a blend of uncapped
- 17 and capped polymer, wherein complete solubilization of the
- 18 copolymer leaves no residual polymer at the site of
- 19 administration and occurs concurrently with the complete release
- 20 of the entrapped agent.
- 21 37. A process of using composition of Claim 1 for human
- 22 administration via parenteral routes, such as intramuscular and
- 24 38. A process of using the composition of Claim 1 for human
- 25 administration via topical route.

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39. A process of using the composition of Claim 1 for human 1

- administration via oral routes.
- 40. A process of using the composition of Claim 1 for human 2
- administration via nasal, transdermal, rectal, and vaginal 3 4
- routes. 5
- 41. A process of using the composition of Claim 1 for human
- administration in the form of an oral or nasal inhalant for the 6 7
- respiratory tract. 8
- 42. A process for preparing controlled release compositions
- characterized by burst-free, sustained, programmable release of 9
- biologically active agents, comprising: Dissolving biodegradable 10 11
- poly(lactide/glycolide), in uncapped form in methylene chloride, 12
- and dissolving a biologically active agent or active core in
- water; adding the aqueous layer to the polymer solution and 13
- emulsifying to provide an innter water-in-oil (w/o) emulsion; 14
- stabilizing the w/o emulsion in a solvent-saturated aqueous phase 15
- containing a oil-in-water (o/w) emulsifier; adding said w/o 16
- emulsion to an external aqueous layer containing oil-in-water 17
- emulsifier to form a ternary emulsion; and stirring the resulting 18
- water-in-oil-in-water (w/o/w) emulsion for sufficient time to 19
- remove said solvent, and rinsing hardened microcapsules with 20
- water and lyophilizing said hardened microcapsules. 21
- The process of Claim 42 wherein a solvent-saturated external 22
- aqueous phase is added to emulsify the inner w/o emulsion prior 23 24
- to addition of the external aqueous layer, to provide 25

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microcapsules of narrow size distribution range between 0.05-1

- 2
- 44. The process of Claim 42 wherein a low temperature of about 3
- 0-4 degree C is provided during preparation of the inner w/o
- emulsion, and a low temperature of about 4-20 degree C is
- provided during preparation of the w/o/w emulsion to provide a 5
- stable emulsion and high encapsulation efficiency. 6
- 45. A process for preparing controlled release compositions 7 8
- characterized by burst-free, sustained compositions characterized 9
- by burst-free, sustained, programmable release of biologically 10
- active agents, comprising: 11
- dissolving biodegradble poly(lactide/glycolide) in end-
- capped form in methylene chloride, and dissolving a biologically 12
- active agent or active core in water; adding the aqueous layer to 13
- the polymer solution and emulsifying to provide an inner water-14
- in-oil emulsion; stabilizing the w/o emulsion in a solvent-15
- saturated aqueous phase containing a oil-in-water (o/w) 16
- emulsifier; adding said w/o emulsion to an external aqueous layer 17
- containing oil-in-water emulsifier to form a ternary emulsion; 18
- and stirring a resulting water-in-oil-water (w/o/w) emulsion for 19 20
- sufficient time to remove said solvent; and rinsing heardened
- microcapsules with water; and lyophilizing said hardened 21 22
- microcapsules. 23
- 46. The process of Claim 42 wherein a 100/0 blend of uncapped
- and end-capped polymer is used to provide release of the artive 24
- core in a continuous and sustained manner without a lag phase. 25 26

- 47. The process of Claim 45 wherein a solvent-saturated external
- aqueous phase is added to emulsify the inner w/o emulsion prior 1
- to addition of the external aqueous layer, to provide 2
- microcapsules of narrow size distribution range between 0.05-3 4
- 5
- The process of Claim 45 wherein a low temperature of about 500um. 6
- 0-4 degree C is provided during preparation of the inner w/o 7
- emulsion, and a low temperature of about 4-20 degree C is
- provided during preparation of the w/o/w emulsion to provide a 8 9
- stable emulsion and high encapsulation efficiency.
- 49. A method for the protection against infection of a mammal by 11
- pathogenic organisms comprising administering orally to said 12
- mammal an immunogenic amount of an immunostimulating composition 13
- consisting essentially of an antigenic synthetic peptide
- encapsulated within a poly(lactide/galactide) matrix. 14
- The method of Claim 49 wherein the poly(lactide/glycolide) 15
- is a blend of uncapped and end-capped forms, in ratios ranging 16 17
- from 100/0 to 1/99.
- The method of Claim 49 wherein the poly(lactide/glycolide) 18
- is a blend of uncapped and end-capped forms in ratios ranging 19 20
- from 90/10 to 40/60. 21
- The method of Claim 49 wherein the infection is a backerial 22
- The method of Claim 49 where the synthetic peptide cantains infection. 23
- an epitope selected from the group consisting of CFA/I piles 24
- protein T-cell epitopes, B-cell epitopes or mixtures thereof. 25 26

- 54. The method of Claim 49 wherein the infection is a viral 1
- infection. 2
- 55. The method of Claim 49 wherein the infection is parasitic 3
- 56. The method of Claim 49 wherein the infection is a fungal
- infection.
- 57. The method of Claim 52 wherein the bacterial infection is 7
- caused by a bacteria selected from the group consisting 8
- essentially of Salmonella typhi, Shigella Sonnei, Shigella 9
- Flexneri, Shigella dysenteriae, Shigella boydii, Escheria coli, 10
- Vibrio cholera, Group D-2, Group E, Group G, Group I, Group 1, 11
- Listeria, Erysipelothrix, Mycobacterium, Aerobic pathogenic 12
- Actinomycetales, Enterobacteriaceae, Vibrio, aeromonas,
- Plesiomonas, Helicobacter, W. succinogenes, Acineto bacter spp., 13 14
- Foavobacterium, Pseudomonas, Legionella, Brucella, Haemophilus,
- Bordetalla, Mycoplasmas, Gardnerella, Streptobacillus, Spirillum, 15
- Calvmmatobacterium, Clostridium, Treponema, Borrelia, Leptospira, 16
- Anaerobic Gram-negative Bacteria including bacilli and Cocci, 17
- Anaerobic gram-Positive Nonsporeforming Bacilli and Cocci, 18
- yersinia, staphylococcus, clostridium, Enteroccus, Streptoccus, 19
- Aerococcus, Planococcus, Stomatococcus, Micrococcus, Lactoccus, 20 21
- Germella, Pediococcus, Leuconostoc, Bacillus, Neisseria,
- Branhamella, Corvne bacterium, campylobacter, Arcanobacterium 22
- haemolyticum, Rhodococcus spp. Rhodococcus, Group A-4. 23 24
- 58. The method in accordance with Claim 49 comprising 25
- administering orally to said mammal an immunogenic amount of a 26

- pharmaceutical composition consisting essentially of an antigenic 1
- synthetic peptide in the amount of .1 to 1%. 2
- 59. A vaccine for the immunization of a mammal against infection 3
- caused by pathogenic organisms prepared from the composition of
- Claim 1.
- 60. The vaccine according to Claim 59 wherein the polymeric 6
- substance is poly(DL-lactide-co-glycolide).
- 61. The vaccine according to Claim 60 wherein the relative ratio 7 8
- between the lactide and glycolide (L/G) component is within the 9
- range of 40/60 to 0/100. 10
- 62. The vaccine according to Claim 61 wherein the relative ratio 11
- between the amount of lactide and glycolide component is within 12
- the range of 90/10 to 40/60. 13
- 63. A vaccine according to Claim 62 wherein the pathogenic 14
- organisms are bacterial. 15
- 64. A vaccine according to Claim 62 wherein the pathogenic 16
- organisms are viral.
- 65. A vaccine according to Claim 62 wherein the pathogenic 17 18
- organisms are fungal. 19
- 66. A vaccine according to Claim 62 wherein the pathogenic 20
- organisms are parasitic. 21
- 67. The vaccine according to Claim 63 wherein the antigenic 22
- synthetic peptide is selected from the group consisting 23
- essentially of Synthetic Peptides Containing CFA/I Pilus Protein 24
- T-cell Epitopes (Starting Sequence # given) 25
- 4 (Asn-Ile-Thr-Val-Thr-Ala-Ser-Val-Asp-Pro), 26

```
8 (Thr-Ala-Ser-Val-Asp-Pro-Val-Ile-Asp-Leu),
1
              12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
              15(Ile-Asp-Leu-Leu-Gln-Ala-Asp-Gly-Asn-Ala),
2
              20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
3
              26(Pro-Ser-Ala-Val-Lys-Leu-Ala-Tyr-Ser-Pro),
4
5
              72(Leu-Asn-Ser-Thr-Val-Gln-Met-Pro-Ile-Ser),
              78 (Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Gln),
6
             87(Gln-Val-Leu-Ser-Thr-Thr-Ala-Lys-Glu-Phe),
 7
               126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr), and
 8
               133(Gly-Asn-Tyr-Ser-Gly-Val-Val-Ser-Leu-Val), and
 9
 10
 ll mixtures thereof;
  12 Synthetic Peptides Containing CFA/I Pilus Protein B-cell
     (antibody) Eptiopes (Starting Sequence # given)
                3(Lys-Ana-Ile-Thr-Val-Thr-Ala-Ser-Val),
                11(Val-Asp-Pro-Val-Idle-Asp-Leu-Leu-Gln-Ala-Asp),
  14
  15
                22(Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
                32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-Thr-Phe-
  16
  17
                   Glu-Ser-Tyr-Arg-Val),
   18
                 32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe),
   19
                 38(Lys-Thr-Phe-Glu-Ser-Tyr-Arg-Val),
   20
                 66(Pro-Gln-Leu-Thr-Asp-Val-Leu-Asn-Ser),
   21
                 93(Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala),
   22
                 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr),
                  127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
    23
                  124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-
    24
    25
                      Ser), and mixtures thereof; and
    26
```

	Synthetic Peptides Containing CFA/I Pilus Protein T-cell and B-
ì	Synthetic Peptides Containing Cra/1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
2	cell (antibody) Epitopes (Starting Sequence # given)
3	Negatine-Thr-Val-Thr-Aid-Sei-Duz-
4	3(Lys-ABN-110 100) 8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-
5	Ala-Asp),
6	11(Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
7	20 (Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
8	20(Ala-ABP-GL) Here 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-
9	ser), and
10	Ser), and 126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
11	mixtures thereof.
12	claim 67 wherein the Bacteria 10
	consisting essentially or output
13	shigella Flexheri, shassan
14	Figheria coli, Vibrio Chotera,
15	Group G. Group I, Group 1, Listeria, Elisteria
	nathogenic Actinomycevery.
1	Vibrio, aeromonas, Plestyments.
1	reineto bacter SDD., rodiosati
1	9 W. succinogenes, Acthety and Brucella, Haemophilus, Bordetalla, 20 Pseudomonas, Legionella, Brucella, Haemophilus, Spirillum,
2	pseudomonas, Legionella, preptobacillus, Spirillum,
:	Pseudomonas, Legionesia, Streptobacillus, Spirillum, Mycoplasmas, Gardnerella, Streptobacillus, Spirillum, Calymmatobacterium, Clostridium, Treponema, Borrelia, Leptospira,
	22 Calymmatobacterium, Clostricium, Itolyming bacilli and Cocci,
	22 Calymmatobacterium, Closeria including bacilli and Cocci, 23 Anaerobic Gram-negative Bacteria including bacilli and Cocci,
	Nonspore forming bacter
	clostrigium, Entergreen
	26 Aerococcus, Planococcus, Stomatococcus, Micrococcus, Lactoccus,

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Germella, Pediococcus, Leuconostoc, Bacillus, Neisseria,
    Branhamella, Corvne bacterium, campylobacter, Arcanobacterium
1
    haemolyticum, Rhodococcus spp. Rhodococcus, Group A-4.
2
3
          The vaccine according to Claim 67 wherein the antigenic
4
     synthetic peptide is selected from the group consisting
5
     essentially of 4(Asn-Ile-Thr-Val-thr-Ala-Ser-Val-Asp-Pro),
6
 7
     8 (Thr-Ala-Ser-Val-Asp-Pro-Val-Ile-Asp-Leu),
 8
      12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
 9
      15(Ile-Asp-Leu-Leu-Gln-Ala-Asp-Gly-Asn-Ala),
 10
      20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
 11
      26 (Pro-Ser-ala-Val-Lys-Leu-Ala-Tyr-Ser-Pro),
  12
      72(Leu-Asn-Ser-Thr-Val-Gln-Met-Pro-Ile-Ser),
  13
      78 (Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Gln),
  14
      87(Gln-Val-Leu-Ser-Thr-Thr-Ala-Lys-Glu-Phe),
       126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr), and
  15
       133(Gly-Asn-Tyr-Ser-Gly-Val-Val-Ser-Leu-Val), and mixtures
  16
   17
       thereof.
   18
       70. The vaccine according to Claim 69 wherein the antigenic
   19
        synthetic peptide is 4(Asn-Ile-Thr-Val-Thr-Ala-ser-Val-Asp-Pro).
   20
   21
        71. The vaccine according to Claim 69 wherein the antigenic
   22
        synthetic peptide is 8(Thr-ala-ser-Val-Asp-Pro-Val-Ile-asp-Leu).
    23
    24
    25
```

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72. The vaccine according to Claim 69 wherein the antigenic synthetic peptide is 12(Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp). l 2 73. The vaccine according to Claim 69 wherein the antigenic 3 synthetic peptide is 15(Ile-Asp-Leu-Leu-Gln-Ala-Asp-Gly-Asn-Ala). 4 5 The vaccine according to Claim 69 wherein the antigenic 6 synthetic peptide is 20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val). 7 8 The vaccine according to Claim 69 wherein the antigenic 9 synthetic peptide is 26(Pro-Ser-Ala-Val-Lys-Leu-Ala-tyr-Ser-Pro). 10 11 The vaccine according to Claim 69 wherein the antigenic 12 synthetic peptide is 72(Leu-Asn-Ser-Thr-Val-Gln-Met-Pro-Ile-Ser). 13 14 77. The vaccine according to Claim 69 wherein the antigemic 15 synthetic peptide is 78 (Met-Pro-Ile-Ser-Val-Ser-Trp-Gly-Gly-Gln). 16 17 78. The vaccine according to Claim 69 wherein the antigemic 18 synthetic peptide is 87(Gln-Val-Leu-Ser-Thr-thr-Ala-Lys-Glu-Phe). 19 20 79. The vaccine according to claim 69 wherein the antigenic 21 synthetic peptide is 126(Ala-Gly-Thr-Ala-pro-Thr-Ala-Gly-Asn-22 23 24 Tyr). 25

```
80. The vaccine according to Claim 69 wherein the antigenic
    synthetic peptide is 133(Gly-Asn-Tyr-Ser-Gly-Val-Val-Ser-Leu-
1
2
    Val).
     81. The vaccine according to Claim 67 wherein the antigenic
4
     synthetic peptide is selected from the group consisting
 5
     essentially of 3(Lys-Ana-Ile-Thr-Val-Thr-Ala-Ser-Val),
 6
                    11(Val-Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-Asp),
 7
                    22(Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
 8
                    32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-Thr-Phe-
 9
  10
                       Glu-Ser-Tyr-Arg-Val),
                    32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe),
  11
                     38(Lys-Thr-Phe-Glu-Ser-Tyr-Arg-Val),
  12
                     66(Pro-Gln-Leu-Thr-Asp-Val-Leu-Asn-Ser),
  13
                     93 (Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala),
   14
                     124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr),
   15
                     127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
   16
                      124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-
   17
                         Tyr-Ser), and mixtures thereof.
   18
            The vaccine according to Claim 81 wherein the antigenic
    19
        synthetic peptide is 3(Lys-Ana-Ile-Thr-Val-Thr-Ala-Ser-Val).
    20
             The vaccine according to Claim 81 wherein the antigenic
    21
        synthetic peptide is 11(Val-Asp-Pro-Val-Ile-Asp-Leu-Leu-Gln-Ala-
    22
    23
         84. The vaccine according to Claim 81 wherein the antigmic
         Asp) .
     24
         synthetic peptide is 22(Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val).
     25
     26
```

- The vaccine according to Claim 81 wherein the antigenic synthetic peptide is 32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe-Lys-1 2 Thr-Phe-Glu-Ser-Tyr-Arg-Val). The vaccine according to Claim 81 wherein the antigenic 3 synthetic peptide is 32(Ala-Tyr-Ser-Pro-Ala-Ser-Lys-Thr-Phe). 4 The vaccine according to Claim 81 wherein the antigenic
 - synthetic peptide is 38(Lys-Thr-Phe-Glu-Ser-Tyr-Arg-Val). 6 7
 - The vaccine according to Claim 81 wherein the antigenic 8
 - synthetic peptide is 66(Pro-Gln-Leu-Thr-Asp-Val-Leu-Asn-Ser). 9
 - The vaccine according to Claim 81 wherein the antigenic
 - synthetic peptide is 93 (Ala-Lys-Glu-Phe-Glu-Ala-Ala-Ala). 10
 - 90. The vaccine according to Claim 81 wherein the antigenic 11
 - synthetic peptide is 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr). 12
 - 91. The vaccine according to Claim 82 wherein the antigenic 13 14
 - synthetic peptide is 127(Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-15
 - Ser) . 16
 - The vaccine according to Claim 82 wherein the antigenic 17
 - synthetic peptide is 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-18 19
 - Asn-Tyr-Ser) . 20

- The vaccine according to Claim 67 wherein the antigenic 21 22
- synthetic peptide is selected from the group consisting 23
- essentially of 3(Lys-Asn-Ile-Thr-Val-Thr-Ala-Ser-Bal-Asp-Pro),
- 8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-Ala-Asp), 24 25
- 11(Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-Ala-Asp), 26

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20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val),
    124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and
    126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-Ser), and mixtures
3
     thereof.
4
     94. The vaccine according to Claim 93 wherein the antigenic
5
     synthetic peptide is 3(Lys-Asn-Ile-Thr-Val-Thr-Ala-Ser-Bal-Asp-
6
7
     Pro).
8
      95. The vaccine according to Claim 93 wherein the antigenic
 9
      synthetic peptide is 8(Thr-Ala-Ser-Bal-Asp-Pro-Bal-Ile-Asp-Leu-
 10
 11
      Leu-Gln-Ala-Asp).
  12
      96. The vaccine according to Claim 93 wherein the antigenic
  13
      synthetic peptide is 11(Bal-Asp-Pro-Bal-Ile-Asp-Leu-Leu-Gln-ala-
  14
  15
       Asp) .
  16
       97. The vaccine according to Claim 93 wherein the antigenic
   17
       synthetic peptide is 20(Ala-Asp-Gly-Asn-Ala-Leu-Pro-Ser-Ala-Val).
   18
   19
       98. The vaccine according to Claim 93 wherein the antigenic
   20
       synthetic peptide is 124(Lys-Thr-Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-
   21
        Asn-Tyr-Ser) .
    23
    24
    25
    26
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The vaccine according to Claim 93 wherein the antigenic synthetic peptide is 126(Ala-Gly-Thr-Ala-Pro-Thr-Ala-Gly-Asn-Tyr-1 2 Ser). 3 The method of Claim 54, wherein the viral infection is 5 caused by a virus selected from the group consisting essentially of hepatitis A, hepatitis B, hepatitis C, Varicella-Zoster virus, 6 7 Epstein-Barr virus, Rotaviruses, polio virus, human immunodeficiency virus (HIV), herpes simplex virus type 1, human 8 retroviruses, herpes simplex virus type 2, Ebola virus, 10 cytomegalo viruses, Herpes Simplex viruses, Human 11 cytomegalovirus, Varicella-Zoster Virus, Epstein-Barr Virus, Poxvirus, Influenza viruses, Parainfluenza viruses, Respiratory 12 13 Syncytial virus, Rhinoviruses, Coronaviruses, Adenoviruses, Measles virus, Mumps virus, Robella Virus, Human Parvoviruses, 14 Arboviruses, Rabies virus, Enteroviruses, reoviruses, Viruses 15 16 Causing gastroenteritis Hepatitis Viruses, Filoviruses, 17 Arenaaviruses, Papillomaviruses, Polyomaviruses, Human 18 Immunodeficiency viruses, Human Retroviruses, and Spongiform 19 Encephalopathies. 20 21 101. The method in accordance with Claim 49 comprising 22 administering orally to said mammal an immunogenic amount of a pharmaceutical composition consisting essentially of an antigen 23 24 in the amount of .1 to 1%.

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102. A vaccine for the immunization of a mammal against 1 infection by pathogenic organisms consisting essentially of an 2 antigen in the amount of 0.1 to 1% encapsulated within a 3 biodegradable-biocompatible polymeric poly(DL-lactide-co-4 glycolide) matrix wherein the polymer is end-capped or a blend of 5 uncapped and end-capped polymers. 6 7 The vaccine according to Claim 100 wherein the polymer is a 8 blend of end-capped and uncapped polymers. 9 10 104. The vaccine according to Claim 103 wherein the relative 11 ratio between the lactide and glycolide component is within the 12 range of 90/10 to 40/60. 13 14 The vaccine according to Claim 103 wherein the relative 15 ratio between the amount of lactide and glycolide component is 16 within the range of 48/52 to 52/48. 17 18 The vaccine according to Claim 102 wherein the antigen is a 19 bacteria or derivatives thereof. 20 21 107. The vaccine according to Claim 103 wherein the antigen is a 22 virus or derivatives thereof. 23 . 24

25 108. The vaccine according to Claim 103 wherein the antigens is 26 a parasite or derivative thereof.

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1 109. The vaccine according to Claim 103 wherein the antigen is a

² fungus or derivative thereof.

3

- 4 110. The vaccine according to Claim 106 wherein the bacteria is
- selected from the group consisting essentially of Salmonella
- typhi, Shigella Sonnei, Shigella Flexneri, Shigella dysenteriae,
- Shigella boydii, Escheria coli, Vibrio cholera, Group D-2, Group
- E, Group G, Group I, Group 1, Listeria, Erysipelothrix,
- 9 Mycobacterium, Aerobic pathogenic Actinomycetales,
- 10 Enterobacteriaceae, Vibrio, aeromonas, Plesiomonas, Helicobacter,
- W. succinogenes, Acineto bacter spp., Foavobacterium,
- 12 Pseudomonas, Legionella, Brucella, Haemophilus, Bordetalla,
- Mycoplasmas, Gardnerella, Streptobacillus, Spirillum,
- Calymmatobacterium, Clostridium, Treponema, Borrelia, Leptospira.
- Anaerobic Gram-negative Bacteria including bacilli and Cocci,
- Anaerobic gram-Positive Nonsporeforming Bacilli and Cocci,
- yersinia, staphylococcus, clostridium, Enteroccus, Streptoccus,
- 18 Aerococcus, Planococcus, Stomatococcus, Micrococcus, Lactoccus,
- 19 Germella, Pediococcus, Leuconostoc, Bacillus, Neisseria,
- 20 Branhamella, Corvne bacterium, campylobacter, Arcanobacterium
- haemolyticum, Rhodococcus spp. Rhodococcus, Group A-4.

- 23 111. The vaccine of Claim 107 Wherein the virus is selected from
- the group consisting essentially of hepatitis A, hepatitis B,
- hepatitis C, Varicella-Zoster virus, Epstein-Barr virus,
- Rotaviruses, polio virus, human immunodeficiency virus (HIV),

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herpes simplex virus type 1, human retroviruses, herpes simplex virus type 2, Ebola virus, cytomegalo viruses, Herpes Simplex 1 viruses, Human cytomegalovirus, Varicella-Zoster Virus, Epstein-2 Barr Virus, Poxvirus, Influenza viruses, Parainfluenza viruses, 3 Respiratory Syncytial virus, Rhinoviruses, Coronaviruses, 4 Adenoviruses, Measles virus, Mumps virus, Robella Virus, Human 5 6 Parvoviruses, Arboviruses, Rabies virus, Enteroviruses, reoviruses, Viruses Causing gastroenteritis Hepatitis Viruses, 7 Filoviruses, Arenaaviruses, Papillomaviruses, Polyomaviruses, 8 Human Immunodeficiency viruses, Human Retroviruses, and 9 10 Spongiform Encephalopathies. 11 112. An immunostimulating composition comprising encapsulating-12 microspheres, which may contain a pharmaceutically-acceptable 13 adjuvant, wherein said microspheres having a diameter between 1 14 nanogram (ng) to 10 microns (um) are comprised of (a) a 15 biodegradable-biocompatible poly (DL-lactide-co-glycolide) as the 16 bulk matrix, wherein the copolymer (lactide to glycolide L/G) 17 ratio for uncapped and end-capped polymer is 0/100 to 1/99 and 18 (b) an immunogenic substance comprising a bacteria, virus, 19 fungus, parasite, or derivative thereof, that serves to elicit 20 21 the production of antibodies in animal subjects. 22 113. An immunostimulating composition according to Claim 112 23 wherein the amount of said immunogenic substance is within the 26 range of 0.1 to 1.5% based on the volume of said bulk matrix.

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114. An immunostimulating composition according to Claim 10 wherein the immunogenic substance comprises Colony Factor Antigen 1 (CFA/II), hepatitis B surface antigen (HBsAg), a mixture thereof 2 3 physiologically similar antigen. 115. An immunostimulating composition according to Claim 113 5 wherein the relative ratio between the lactide and glycolide 6 7 component is within the range of 48/52 to 52/48. 8 116. An immunostimulating composition according to Claim 113 9 wherein the size of more than 50% of said microspheres is between 10 11 5 to 10 um in diameter by volume. 12 117. An immunostimulating composition according to Claim 113 13 wherein the immunogenic substance is the synthetic peptide representing the peptide fragment beginning with the amino acid residue 63 through 78 of Pilus Protein CS3, said residue having the amino acid sequence, 63(Ser-Lys-Asn-Gly-Thr-Val-Thr-Try-Ala-17 18 His-Glu-Thr-Asn-Asn-Ser-Ala). 19 118. A vaccine comprising an immunostimulating composition of 20 Claim 113 and a sterile, pharmaceutically-acceptable carrier 21 22 therefor. 23 24 25

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119. A vaccine comprising an immunostimulating composition of Claim 118 wherein said immunogenic substance is Colony Factor 1 2 Antigen (CFA/II). 3 120. A vaccine comprising an immunostimulating composition of 4 Claim 119 wherein said immunogenic substance is hepatitis B 5 6 surface antigen (HBsAg). 121. A method for the vaccination against bacterial infection 8 comprising administering to a human, an antibactericidally 9 10 effective amount of a composition of Claim 118. 11 122. A method according to Claim 121 wherein the bacterial 12 infection is caused by a bacteria selected from the group 13 consisting essentially of Salmonella typhi, Shigella Sonnei, 14 Shigella Flexneri, Shigella dysenteriae, Shigella boydii, 15 Escheria coli, Vibrio cholera, Group D-2, Group E, Group G, Group 16 I, Group 1, Listeria, Erysipelothrix, Mycobacterium, Aerobic 17 pathogenic Actinomycetales, Enterobacteriaceae, Vibrio, 18 aeromonas, Plesiomonas, Helicobacter, W. succinogenes, Acineto 19 bacter spp., Foavobacterium, Pseudomonas, Legionella, Brucella, 20 21 Haemophilus, Bordetalla, Mycoplasmas, Gardnerella, Streptobacillus, Spirillum, Calymmatobacterium, Clostridium, 22 Treponema, Borrelia, Leptospira, Anaerobic Gram-negative Bacteria 23 including bacilli and Cocci, Anaerobic gram-Positive 24 Nonsporeforming Bacilli and Cocci, versinia, staphylococcus, 25

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clostridium, Enteroccus, Streptoccus, Aerococcus, Planococcus, ١. Stomatococcus, Micrococcus, Lactoccus, Germella, Pediococcus, Leuconostoc, Bacillus, Neisseria, Branhamella, Corvne bacterium, 2 campylobacter, Arcanobacterium haemolyticum, Rhodococcus spp.. 3 4 Rhodococcus, Group A-4. 5 A method for the vaccination against viral infection 6 comprising administering to a human an antivirally effective 7 8 amount of a composition of Claim 108. 9 A diagnostic assay for bacterial infections comprising a 10 11 composition of Claim 7. 12 125. A method of preparing an immunotherapeutic agent against 13 infections caused by a bacteria comprising the steps of (1) 14 immunizing a plasma donor with a vaccine according to Claim 52 15 such that a hyperimmune globulin is produced which contains 16 antibodies directed against the bacteria; (2) separating the 17 hyperimmune globulin and (3) purifying the hyperimmune globulin. 18 19 126. A method preparing an immunotherapeutic agent against 20 infections caused by a virus comprising the step of immunizing a 21 plasma donor with a vaccine according to Claim 126 such that 22 hyperimmune globulin is produced which contains antibodies 23 24 directed against the hepatitis B virus. 25 26

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127. An immunotherapy method comprising the step of administering to a subject an immunostimulatory amount of 1 hyperimmune globulin prepared according to Claim 125. 2 3 128. An immunotherapy method comprising the step of 4 administering to a subject an immunostimulatory amount of 5 hyperimmune globulin prepared according to Claim 125. 6 7 129. A method for the protection against infection of a subject 8 by enteropathogenic organisms or hepatitis B virus comprising 9 administering to said subject an immunogenic amount of an 10 11 immunostimulating composition of Claim 112. 12 130. A method according to Claim 127 wherein the 13 immunostimulating composition is administered orally. 14 15 131. A method according to Claim 127 wherein the 16 immunostimulating composition is administered parenterally. 17 18 132. A method according to Claim 127 wherein the 19 immunostimulating composition is administered in four separate 20 21 doses on day 0, day 7, day 14, and day 28. 22 133. A method according to Claim 114 wherein the immunogenic 23 substance is the synthetic peptide representing the peptide 24 fragment beginning with the amino acid residue 63 through 78 of 25 26

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Pilus Protein CS3 said residue having the amino acid sequence 63 (Ser-Lys-Asn-Gly-Thr-Val-Thr-Try-ala-His-Glu-thr-asn-Asn-Ser-1 2 Ala). 3 134. A method for the protection against or therapeutic 4 treatment of bacterial infection in the soft tissue or bone of a 5 mammal comprising administering locally to said mammal a 6 bactericidally-effective amount of a composition of Claim 2, 7 wherein the active material is an antibiotic which is controlled 8 release within a period of about 1 to 100 days. 10 The method according to Claim 134 wherein the biodegradable 11 poly(DL-lactide-co-glycolide) is a blend of uncapped and end-12 capped forms having a relative ratio between the amount of 13 lactide and glycolide component within the range of 100/0 to 14 15 1/99. 16 136. A method according to Claim 135 wherein the bacterial 17 infection is (1) a subcutaneous infection secondary to 18 contaminated abdominal surgery, (2) an infection surrounding 19 prosthetic devices and vascular grafts, (3) ocular infections, 20 (4) topical skin infections, (5) orthopedic infections, including 21 22 osteomyelitis, and (6) oral infections. 23 24

The method according to Claim 136 wherein the oral 137. 25

infections are pericoronitis or periodontal disease. 26

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138. The method according to Claim 135 wherein the 1 administration is effected prior to infection. 2 3 The method according to Claim 135 wherein the 4 administration is effected subsequent to infection. 5 140. The method according to Claim 135 wherein said animal is a 6 7 8 human. The method according to Claim 135 wherein said animal is a 9 10 11 nonhuman. 142. The method in accordance with Claim 135 comprising applying 12 13 to the soft tissue or bone tissue of said animal a bactericidally-effective amount of a pharmaceutical composition 14 consisting essentially of an antibiotic in the ant, selected from 15 the group consisting of a beta-lactam, aminoglycolide, polymyxin-16 b, Amphotericin B, Aztreonam, cephalosporins, chloramphenicol, 17 fusidans, lincosamides, macrolides, methronidazole, nitro-18 furation, Imipenem/cilastin, quinolones, refampin, polyenes, 19 tetracycline, sulfonamides, trimethoprim, vancomycin, 20 teicoplanin, imidazoles, and erythromycin, encapsulated within a 21 biodegradable poly(DL-lactide-co-glycolide) polymeric matrix, 22 wherein the amount of the lactide and glycolide (L/G) component 23 is within the range of 48/52 to 52/48 based on the weight of said 24 polymeric matrix which is present in the amount of from 40 to 95 25

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percent, resulting in the controlled release of a bacteriacidal amount of the said antibiotic over a period of from 1 to 100 1 2 days. 3 143. The method of Claim 142 wherein the polymeric matrix consists essentially of a poly(DL-lactide-co-glycolide) wherein the relative ratio between the amount of lactide and glycolide 6 7 (L/G) component is within the range of 48/52 to 52/48. 8 144. The method of Claim 142 wherein the bacterial infection is 9 caused by a resistant or non-resistant bacteria selected from the 10 group consisting essentially of Enterobacteriaceae; Klebsiella 11 sp.; Bacteroides sp. Enterococci; Proteus sp.; Streptococcus sp.; 12 13 Staphylococcus sp.; Pseudomonas sp.; Neisseria sp.; 14 Pedptostreptococcus sp.; Fusobacterium sp.; Actinomyces sp.; 15 Mycobacterium sp.; Listeria sp.; Corynebacterium sp.; Proprionibacterium sp.; Actinobacillus sp.; Aerobacter sp.; 16 Borrelia sp.; Campylobacter sp.; cytophaga sp.; Pasteurella sp.; 17 Clostridium sp., Enterobacter aerogenes, Peptococcus sp., Proteus 18 vulgaris, Proteus morganii, Staphylococcus aureus, Streptococcus 19 pyogenes, Actinomyces sp., Campylobacter fetus, and Legionella 20 pneumophila, ampillin-resistant strain of S. aureus, and 21 22 methicillin-resistant strain of S. aureus. 23 24

The method of Claim 142 wherein the antibiotic is selected 25

from the group consisting essentially of a beta-lactam, 26

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aminoglycolide, polymyxin-B, amphotericin B, aztreonam, cephalosporins, chloramphenicol, fusidans, lincosamides, 1 macrolides, methronidazole, nitro-furantoin, Imipenem/cilastin, 2 quinolones, rifampin, polyenes, tetracycline, sulfonamides, 3 trimethoprim, vancomycin, teicoplanin, imidazoles, and 4 5 erythromycin. 146. The method of Claim 145 wherein the beta-lactam is 7 cephalosporin. 9 147. The method of Claim 145 wherein the beta-lactam is 10 11 penicillin. 12 148. The method of Claim 145 wherein the aminoglycolide is 13 14 gentamicin. 15 149. The method of Claim 145 wherein the aminoglycolide is 16 17 amikacin. 18 150. The method of Claim 145 wherein the aminoglycolide is 19 20 tobramycin. 21 151. The method of Claim 145 wherein the aminoglycolide is 22 23 kanamycin. 24

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152. The method of Claim 145 wherein the beta-lactam is an 1 ampicillin. 2 3 The method of Claim 152 wherein the polymeric matrix 4 consists essentially of a poly(DL-lactide-co-glycolide) wherein 5 the relative ratio between the amount of lactide and glycolide 6 (L/G) component is within the range of 48/52 to 58/42. 7 8 154. The method of Claim 152 wherein the ampicillin is present 9 in an amount of from 5 to 60 percent and the amount of polymeric 10 matrix is from 40 to 95 percent. 11 12 155. The process of using the composition of Claim 1 to treat 13 humans in need, thereof, suffering from diseases and/or ailments 14 from the group consisting of: viral infections; bacterial 15 infections; fungal infections; parastic infections and more 16 specific diseases and/or ailments; such as as, aids; alzheimer's 17 dementia; angiogenesis diseases; aphthour ulcers in AIDS 18 patients; asthma; atopic dermatitis; psoriasis; basal cell 19 carcinoma; benign prostatic hypertrophy; blood substitute; blood 20 substitute in surgery patients; blood substitute in trauma 21 patients; breast cancer; breast cancer; cutaneous & metastatic; 22 cachexia in AIDS; campylobacter infection; cancer; pnemonia; 23 sexually transmitted diseases (STDs); cancer; viral dieases; 24 candida albicians in AIDS and cancer; candidiasis in HIV 25 infection; pain in cancer; pancreatic cancer; parkinson's

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disease; peritumoral brain edema; postoperative adhesions (prevent); proliferative diseases; prostate cancer; ragweed ı allergy; renal disease; restenosis; rheumatoid arthritis; 2 rheumatoid arthritis; allergies; rotavirus infection; scalp 3 psoriasis; septic shock; small-cell lung cancer; solid tumors; 4 stroke; thrombosis; type I diabetes; type I diabetes w/kidney 5 transplants; type II diabetes; viseral leishmaniasis; malaria; 6 periodontal or gum disease; cardiac rthythm disorders; central 7 nervous system diseases; central nervous system disorders; 8 cervical dystonia (spasmodic torticollis); choridal 9 neovascularization; chronic hepatitis c, b and a; colitis 10 associated with antibiotics; colorectal cancer; coronary artery 11 thrombosis; cryptosporidiosis in AIDS; cryptosporidium parvum 12 diarrhea in AIDS; cystic fibrosis; cytomegalovirus disease; 13 depression; social phobias; panic disorder; diabetic 14 complications; disabetic eye disease; diarrhea associated with 15 antibiotics; erectile dysfunction; genital herpes; graft-vs host 16 disease in transplant patients; growth hormone deficiency; head 17 and neck cancer; head trauma; stroke; heparin neutralization 18 after cardiac bypass; hepatocellular carcinoma; HIV; HIV 19 infection; huntington's disease; CNS diseases; 20 hypercholesterolemia; hypertension; inflammation; inflammation 21 and angiogensis; inflammation in cardiopulmonary bypass; 22 influenza; migrain head ache; interstitial cystitis; kaposi's 23 sarcoma; kaposi's sarcoma in AIDS; lung cancer; melanoma; 24 molluscum contagiosum in AIDS; multiple sclerosis; neoplastic 25

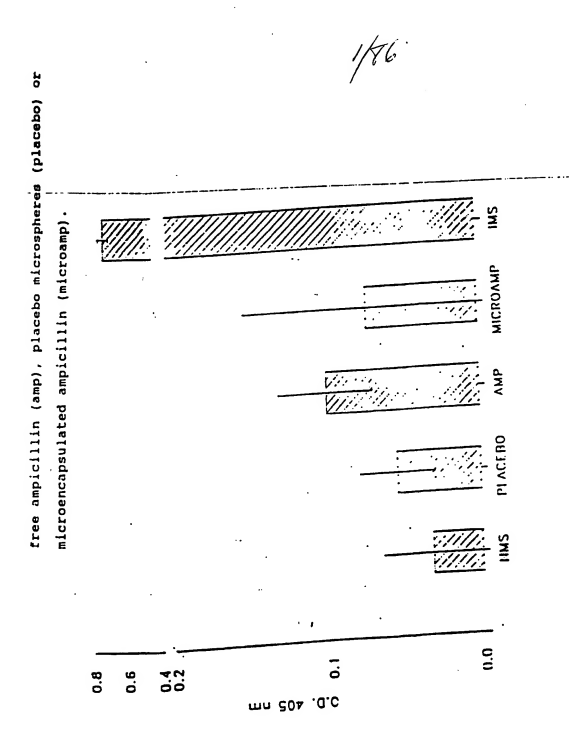
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meningitis from solid tumors; non-small cell lung cancer; organ transplant rejection; osteoarthritis; rheumatoid arthritis; ı osteoporosis; drug addiction; shock; ovarian cancer; Amebiasis; 2 Babesiasis; Chagas' disease (Trypanosoma cruzi); 3 Cryptosporidiosis; Cysticercosis; Pascioliasis; Filariasis; 4 Echinococcosis; Giardiasis; Leishmaniasis; Malaria; 5 Paragonimiasis; Pneumocystosis; Schistosomiasis; Strongylodiasis; Toxocariasis; Toxoplasmosis; Trichinellosis; Trichomoniasis; 7 8 yeast infection; and pain. 9 156. A vaccine for prepared from the composition of Claim 1 to 10 prevent the occurence in humans of diseases and/or ailments 11 comprising viral infections; bacterial infections; fungal 12 infections; parastic infections and more specific diseases and/or 13 ailments; such as as, aids; alzheimer's dementia; angiogenesis 14 diseases; aphthour ulcers in AIDS patients; asthma; atopic 15 dermatitis; psoriasis; basal cell carcinoma; benign prostatic 16 hypertrophy; blood substitute; blood substitute in surgery 17 patients; blood substitute in trauma patients; breast cancer; 18 breast cancer; cutaneous & metastatic; cachexia in AIDS; 19 campylobacter infection; cancer; pnemonia; sexually transmitted 20 diseases (STDs); cancer; viral dieases; candida albicians in AIDS 21 and cancer; candidiasis in HIV infection; pain in cancer; 22 pancreatic cancer; parkinson's disease; peritumoral brain edema; 23 postoperative adhesions (prevent); proliferative diseases; 24 prostate cancer; ragweed allergy; renal disease; restenosis; 25

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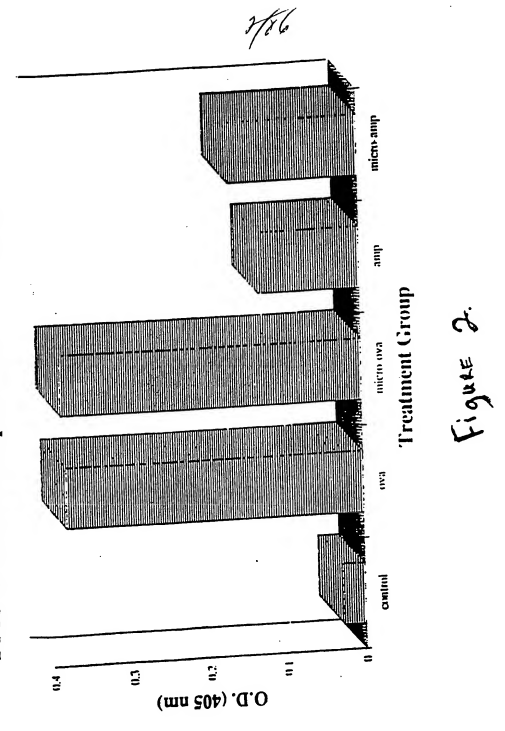
rheumatoid arthritis; rheumatoid arthritis; allergies; rotavirus infection; scalp psoriasis; septic shock; small-cell lung cancer; 1 solid tumors; stroke; thrombosis; type I diabetes; type I 2 diabetes w/kidney transplants; type II diabetes; viseral 3 leishmaniasis; malaria; periodontal or gum disease; cardiac 4 rthythm disorders; central nervous system diseases; central 5 nervous system disorders; cervical dystonia (spasmodic 6 torticollis); choridal neovascularization; chronic hepatitis c, b 7 and a; colitis associated with antibiotics; colorectal cancer; 8 coronary artery thrombosis; cryptosporidiosis in AIDS; 9 cryptosporidium parvum diarrhea in AIDS; cystic fibrosis; 10 cytomegalovirus disease; depression; social phobias; panic 11 disorder; diabetic complications; disabetic eye disease; diarrhea 12 associated with antibiotics; erectile dysfunction; genital 13 herpes; graft-vs host disease in transplant patients; growth 14 hormone deficiency; head and neck cancer; head trauma; stroke; 15 heparin neutralization after cardiac bypass; hepatocellular 16 carcinoma; HIV; HIV infection; huntington's disease; CNS 17 diseases; hypercholesterolemia; hypertension; inflammation; 18 inflammation and angiogensis; inflammation in cardiopulmomary 19 bypass; influenza; migrain head ache; interstitial cystitis; 20 kaposi's sarcoma; kaposi's sarcoma in AIDS; lung cancer; 21 melanoma; molluscum contagiosum in AIDS; multiple sclerosis; 22 neoplastic meningitis from solid tumors; non-small cell lung 23 cancer; organ transplant rejection; osteoarthritis; rheumatoid 24 arthritis; osteoporosis; drug addiction; shock; ovarian cancer; 25 26

chagas' disease (Trypanosoma cruzi);
Amebiasis; Babesiasis; Chagas' disease (Trypanosoma cruzi); Cryptosporidiosis; Cysticercosis; Fascioliasis; Filariasis;
2 Cryptosporidiosis; Cysticercosis, talendaria; 3 Echinococcosis; Giardiasis; Leishmaniasis; Malaria; 3 Echinococcosis; Giardiasis; Leishmaniasis; Strongylodiasis;
Paragonimiasis; Pneumocystosis; Schiberts; Trichomoniasis; Toxocariasis; Toxoplasmosis; Trichinellosis; Trichomoniasis;
5 Toxocariasis; Toxoplasmosis; Illument
6 yeast infection; and pain.
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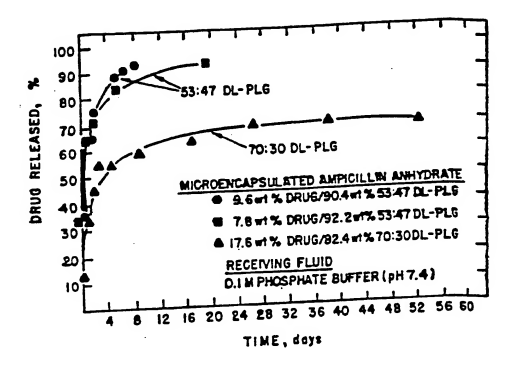


rigure 1

Serum IgG of Guinea Pigs Sensitized with Free vs Microencapsulated OVA and AMP



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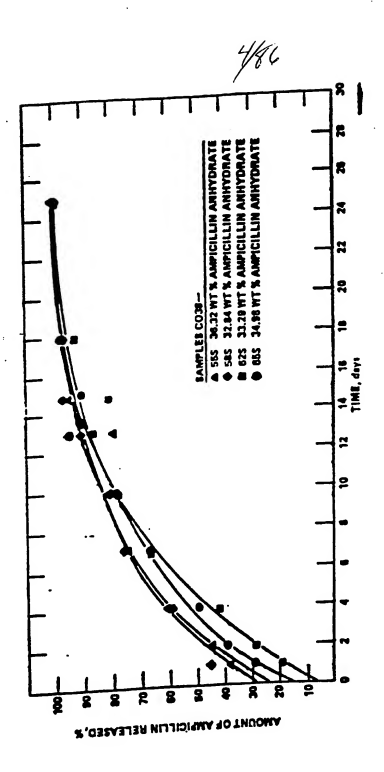


Fig. 7

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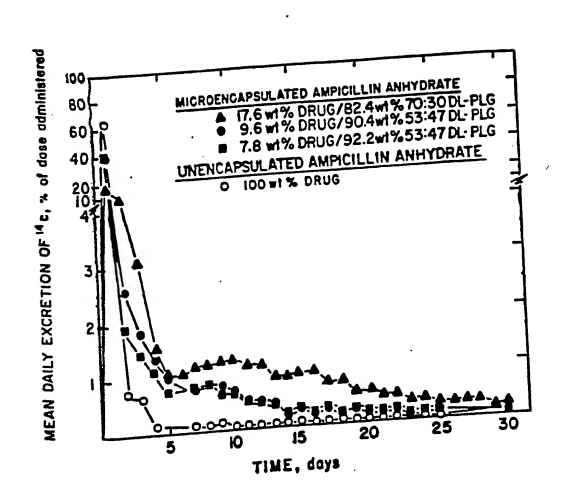


FIG. 5

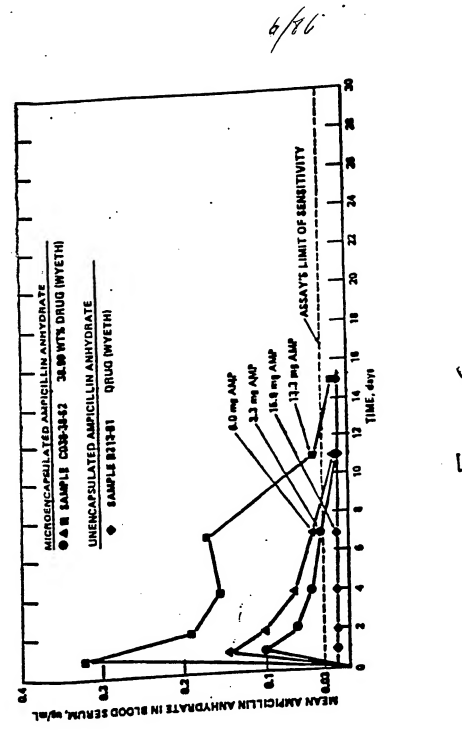


Fig. 6

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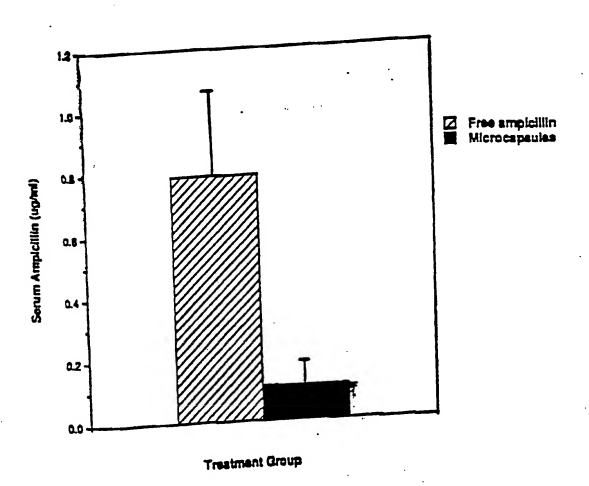


Figure T

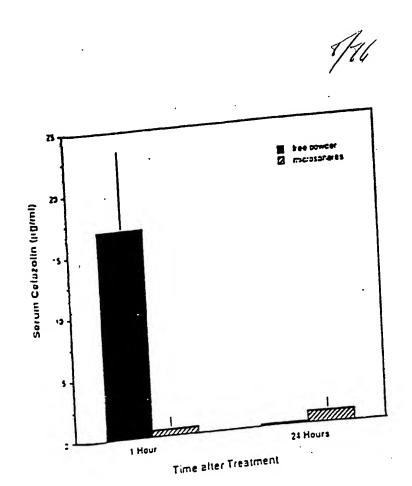
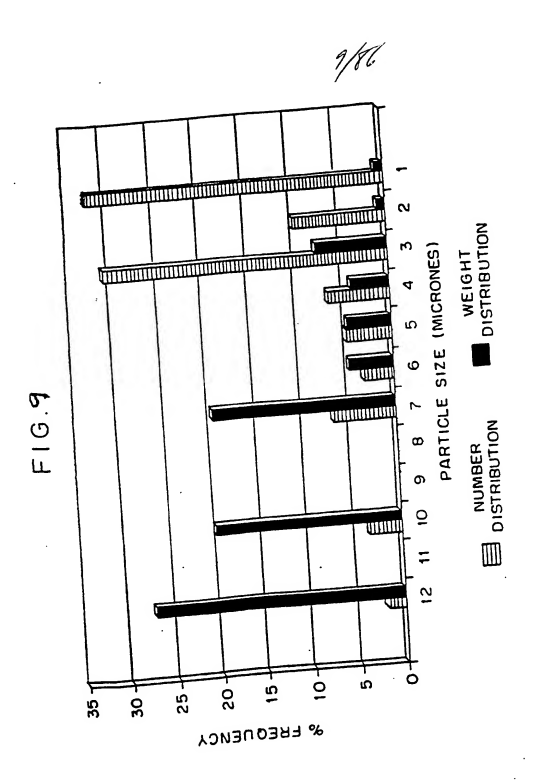


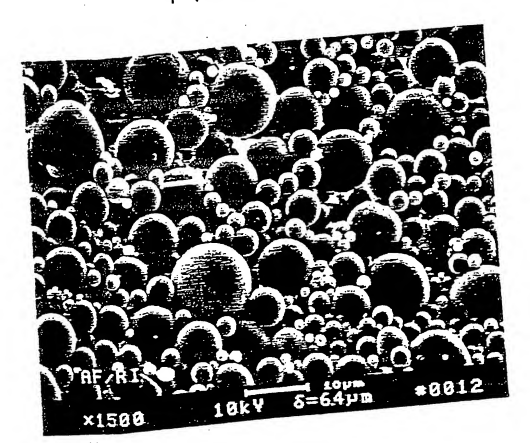
Figure & Serum Cefazolin Levels.

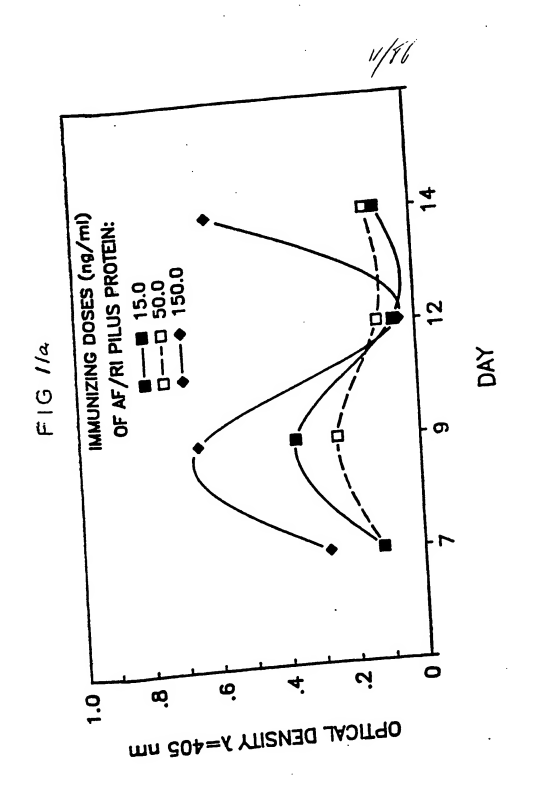


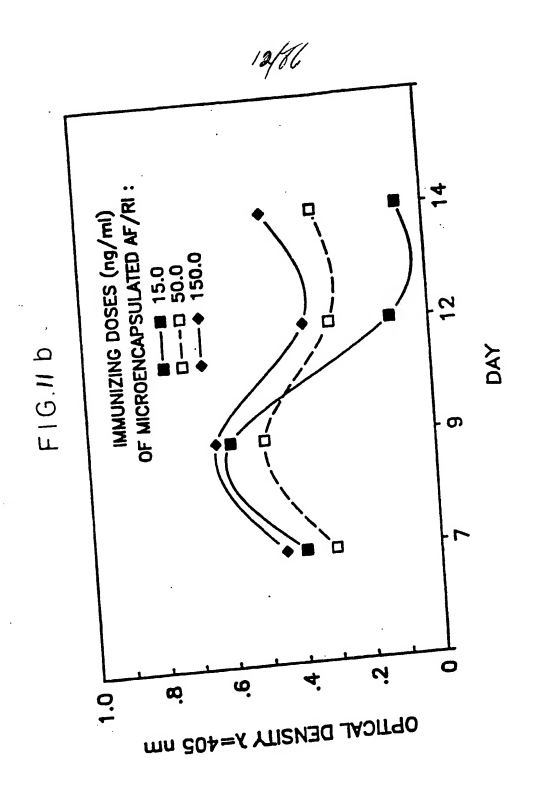
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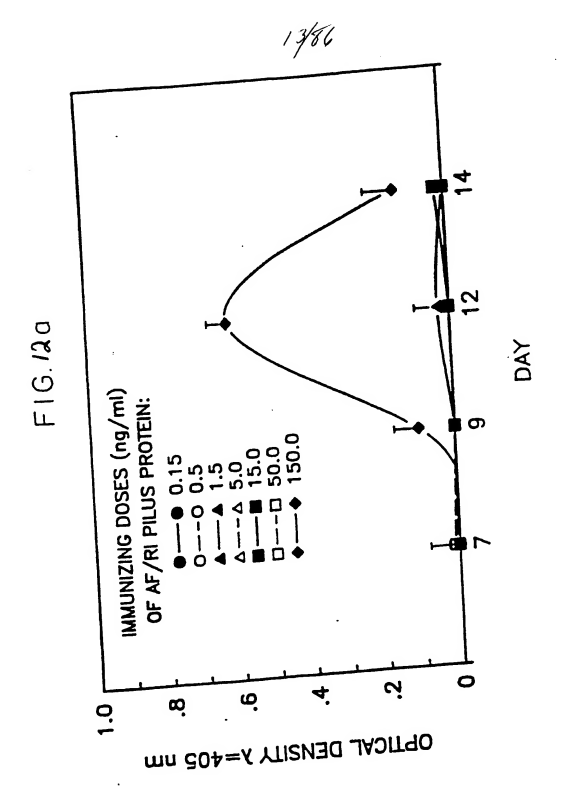
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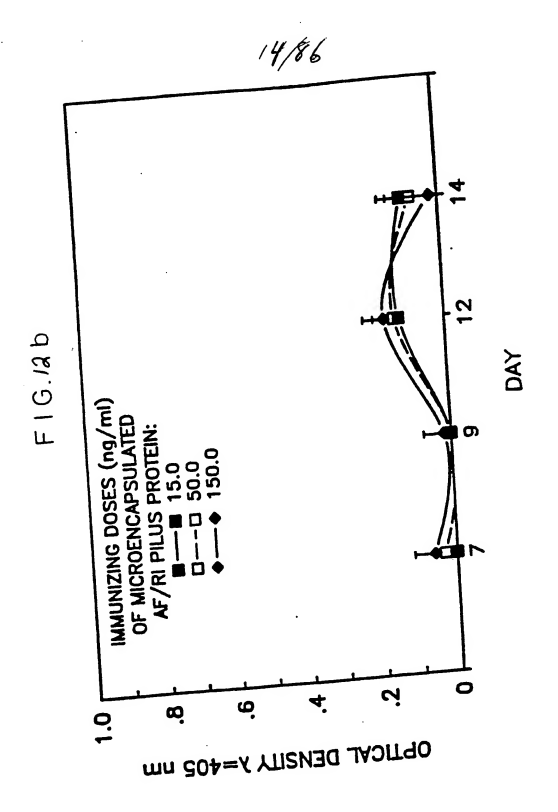
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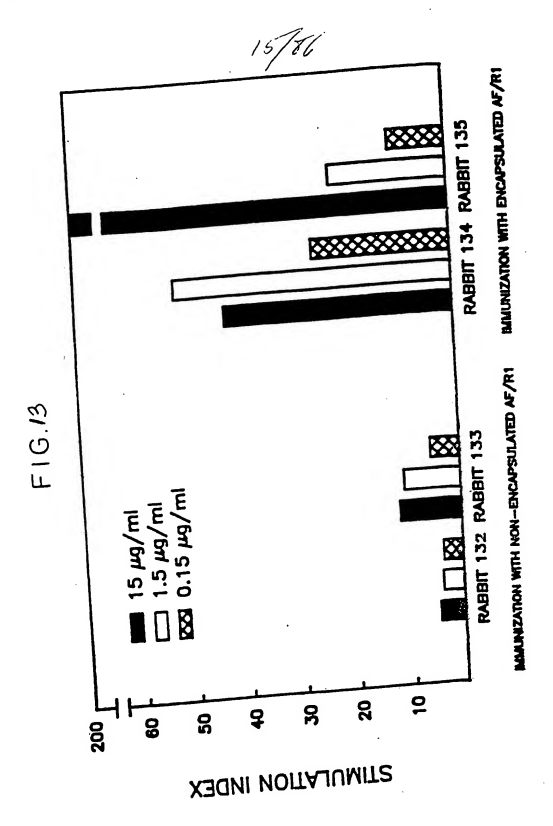


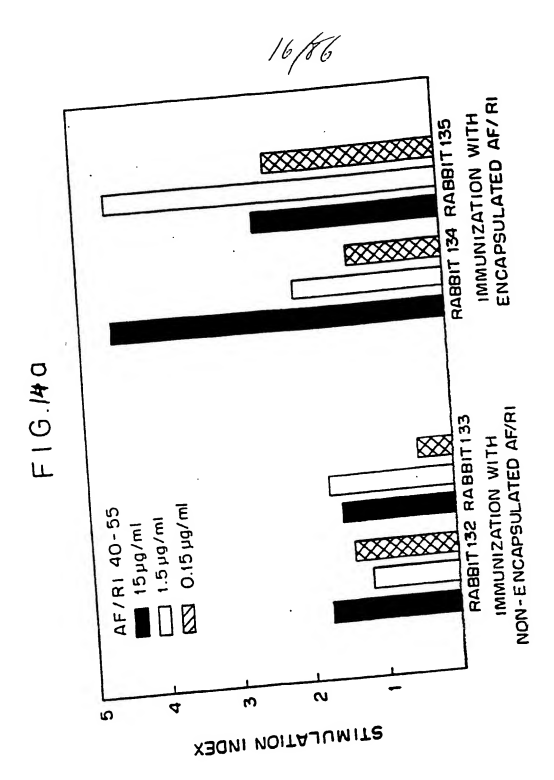


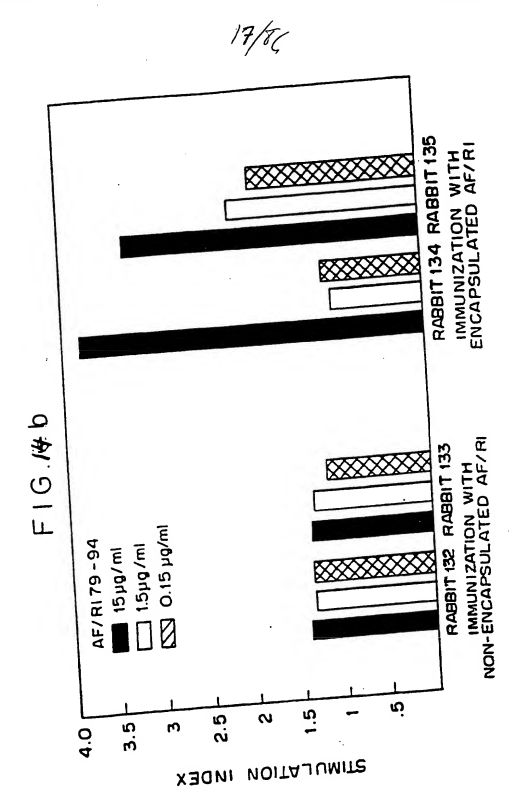


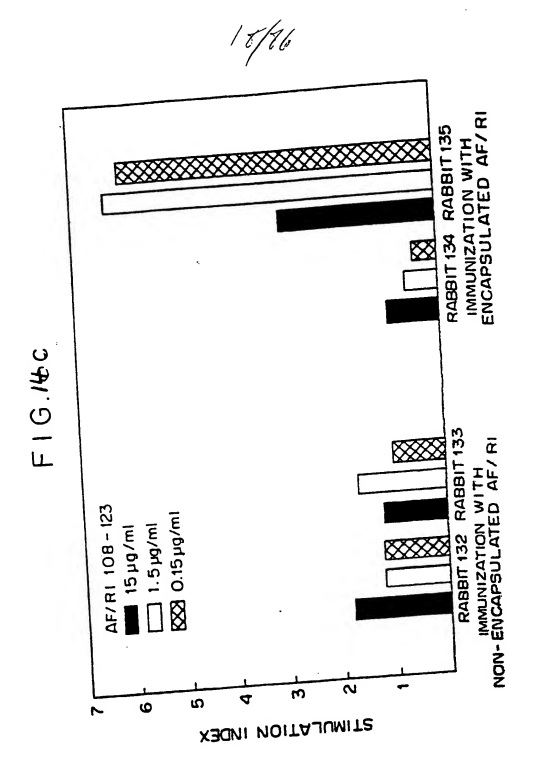


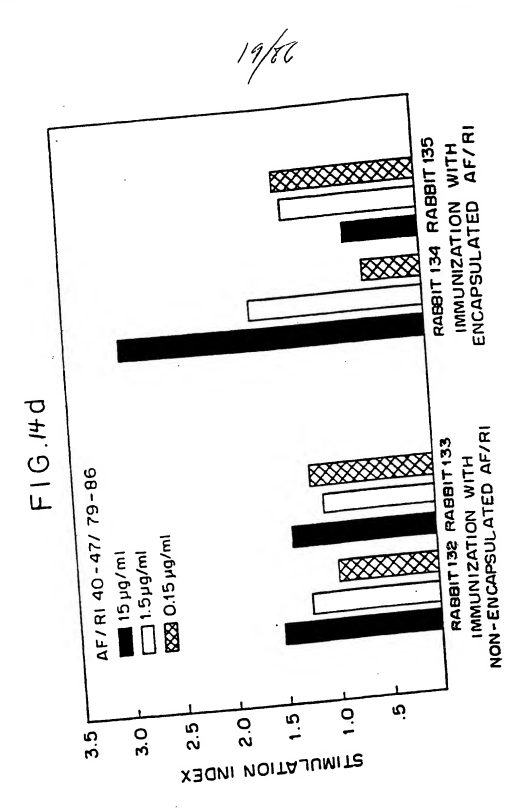


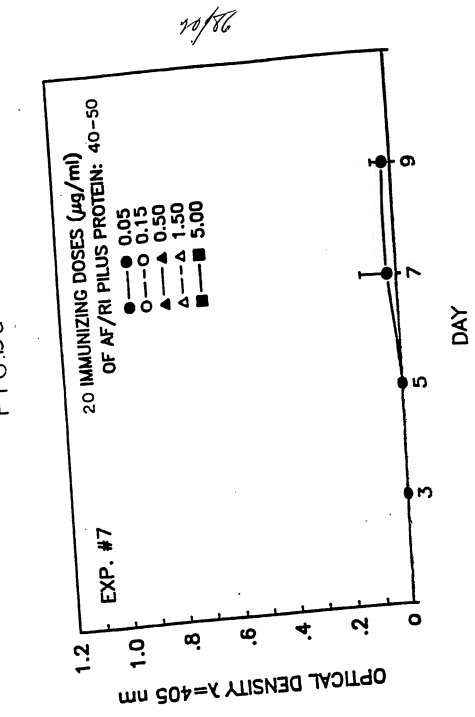




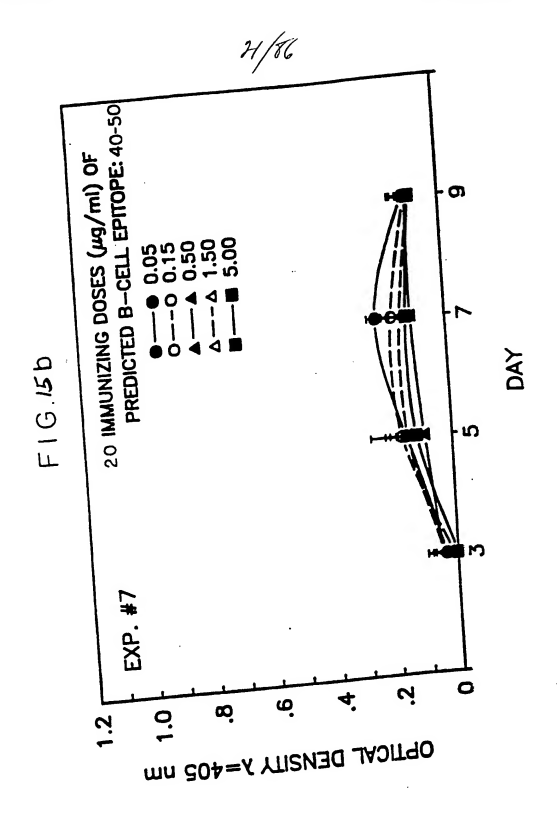


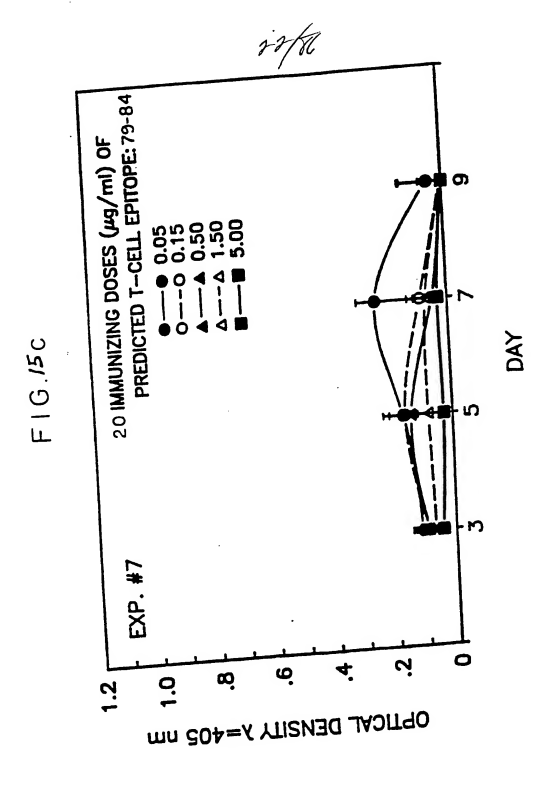


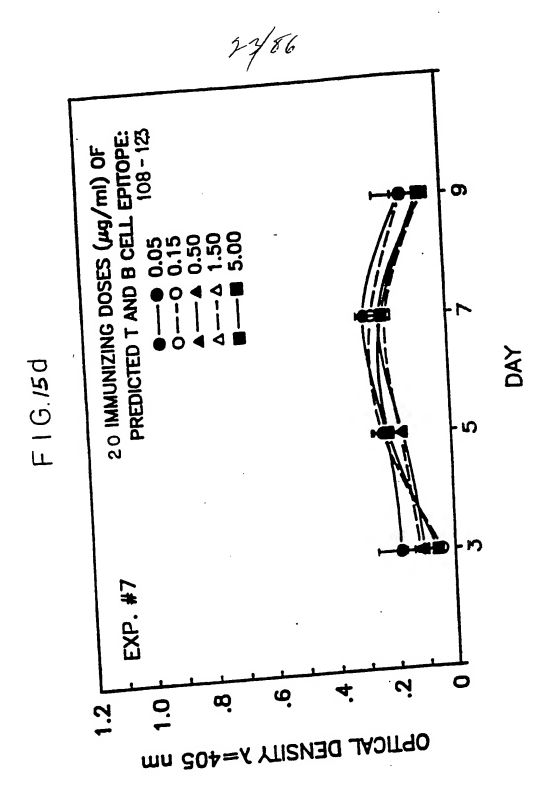


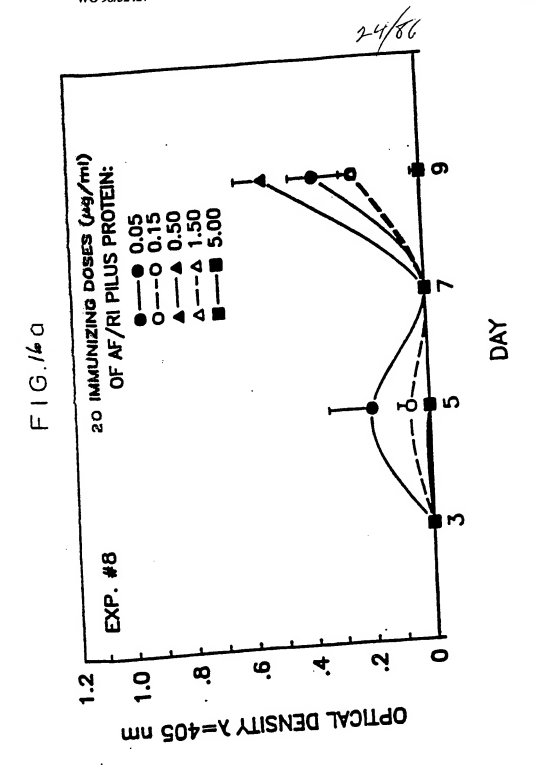


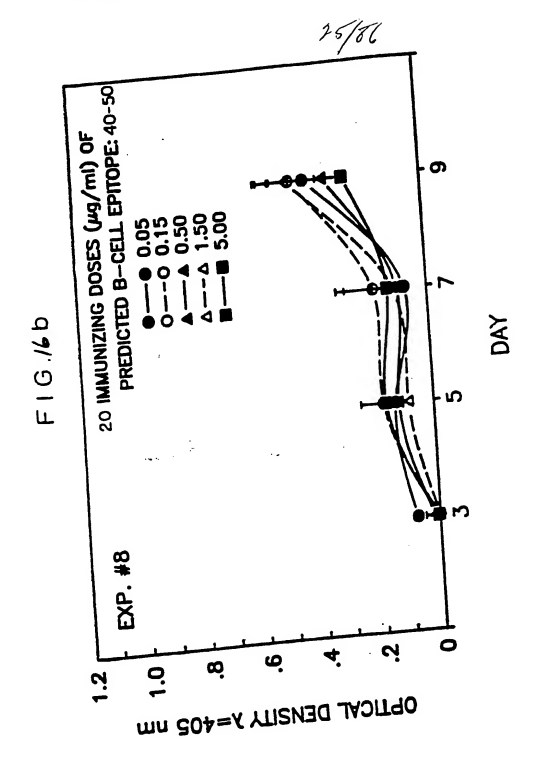
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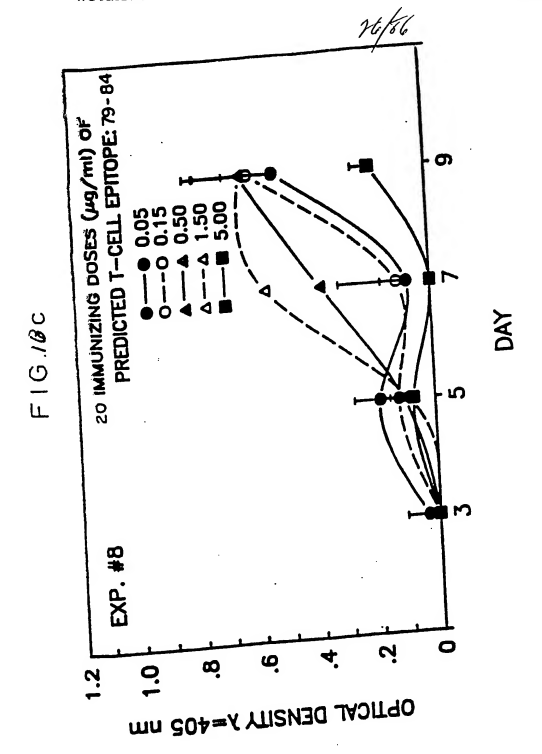


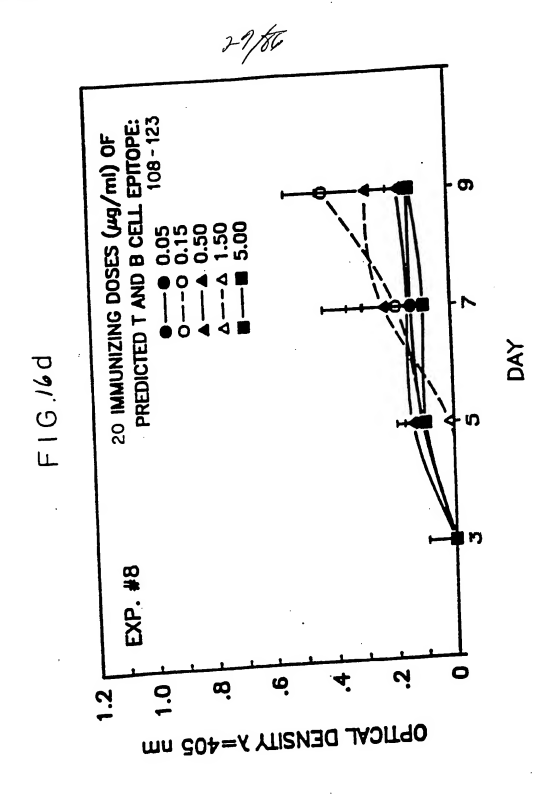


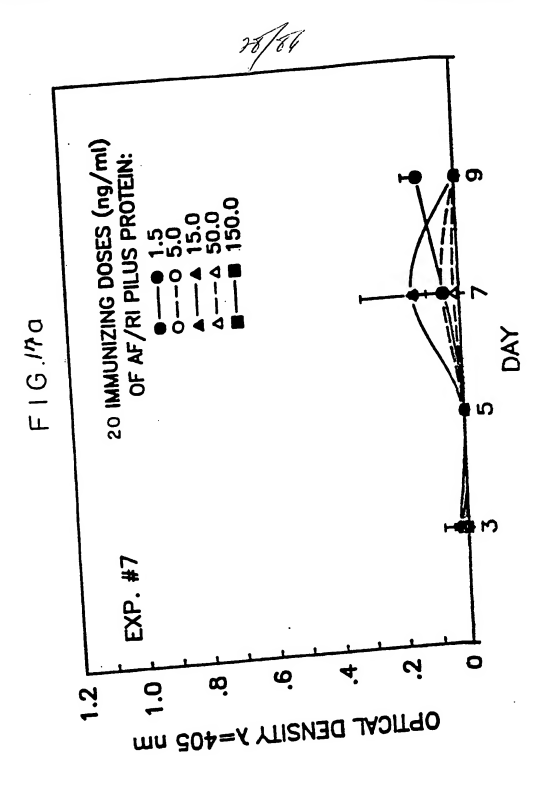


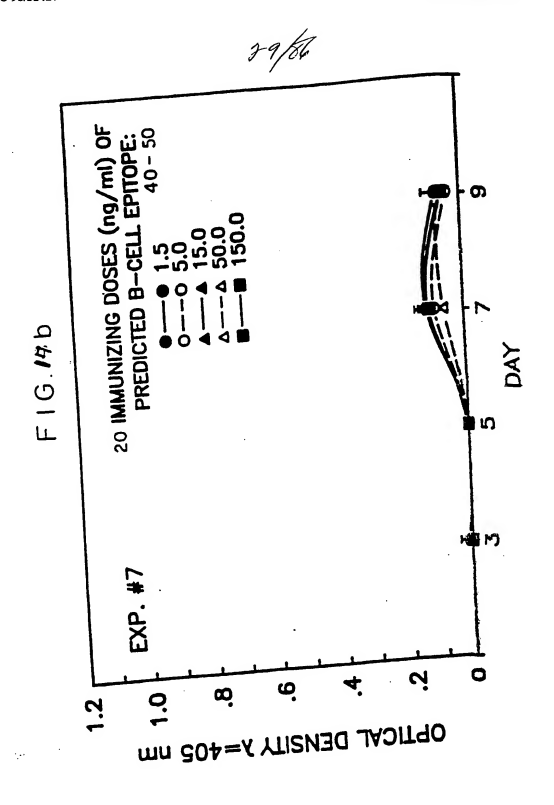


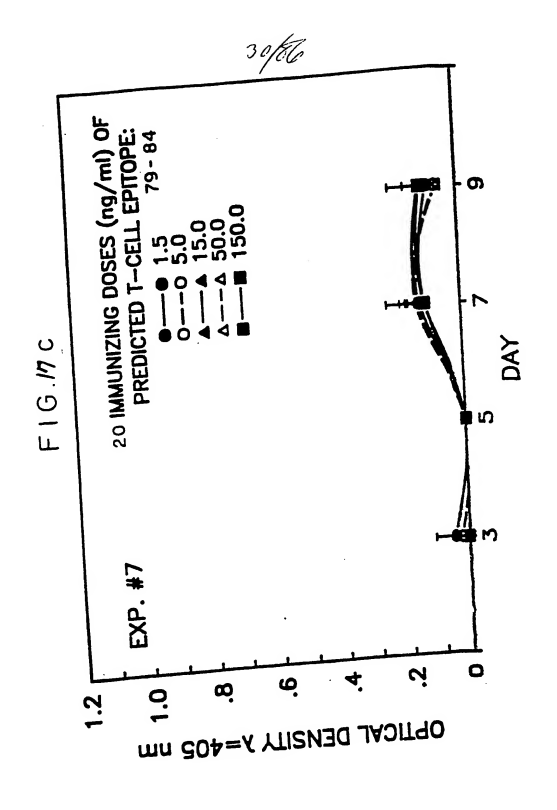


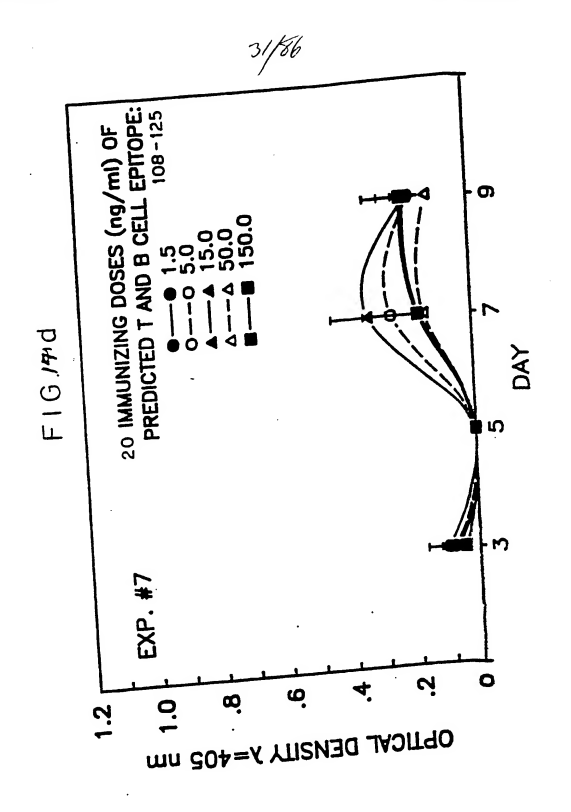


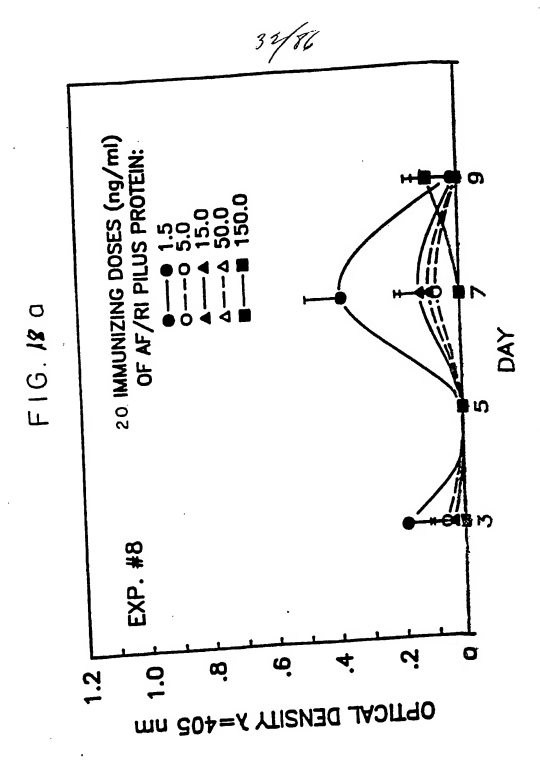


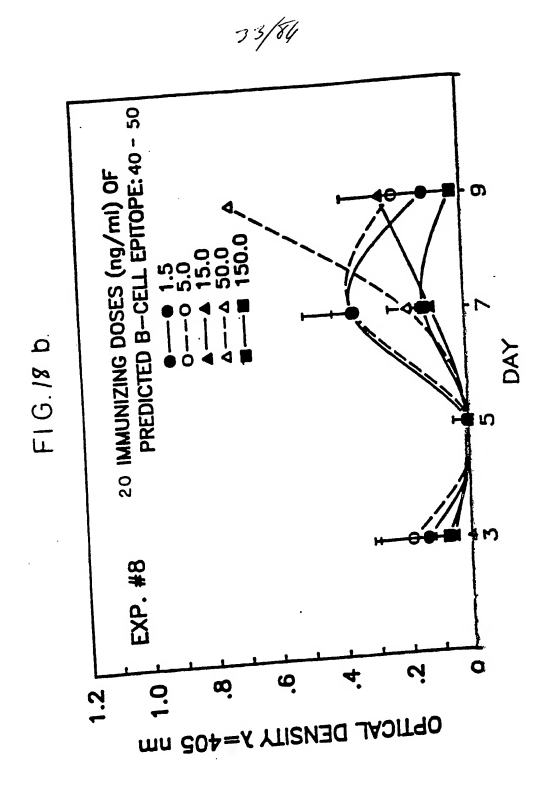


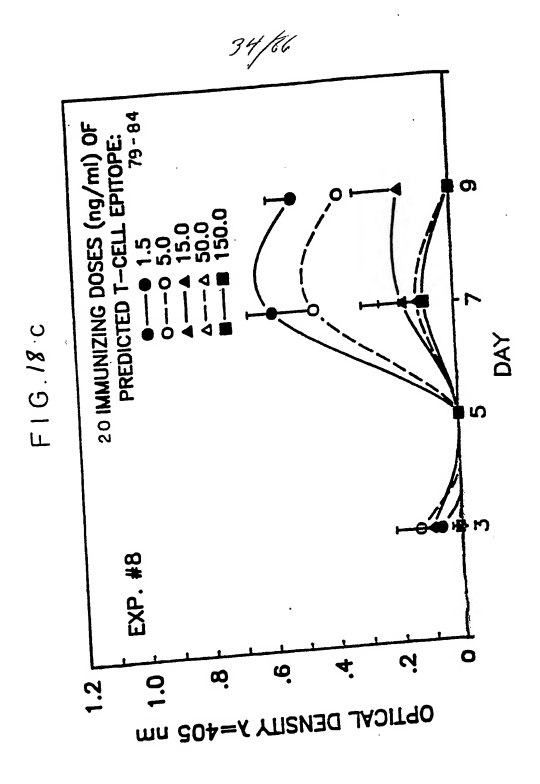




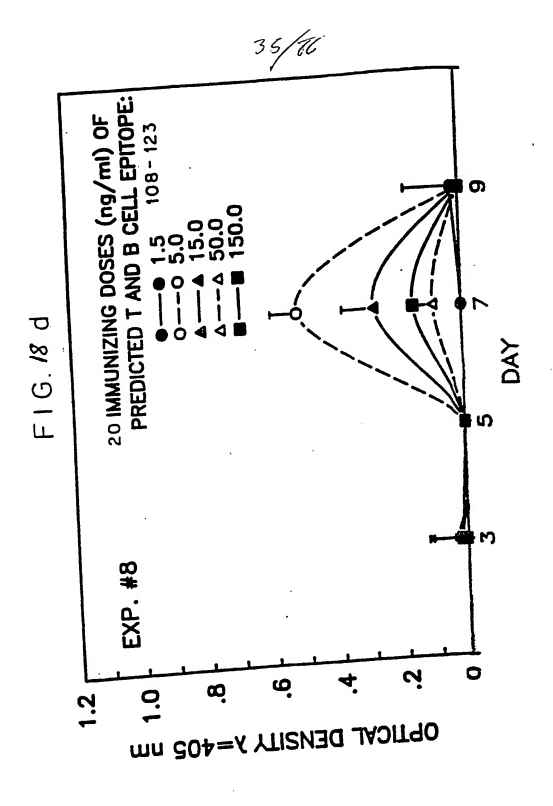


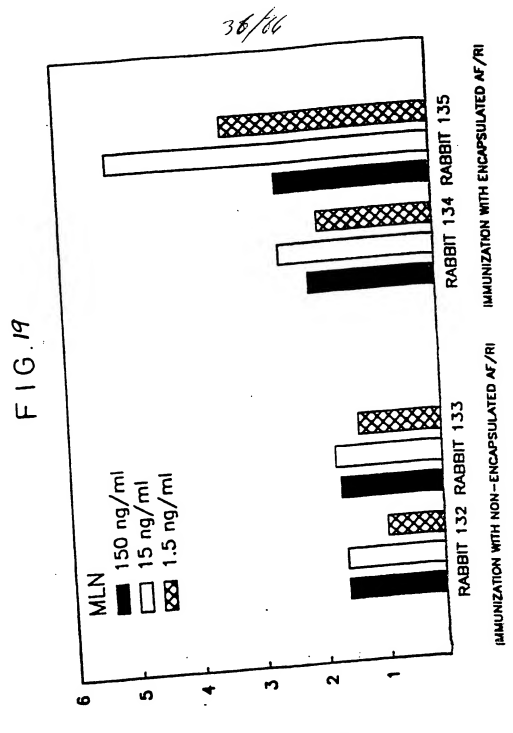




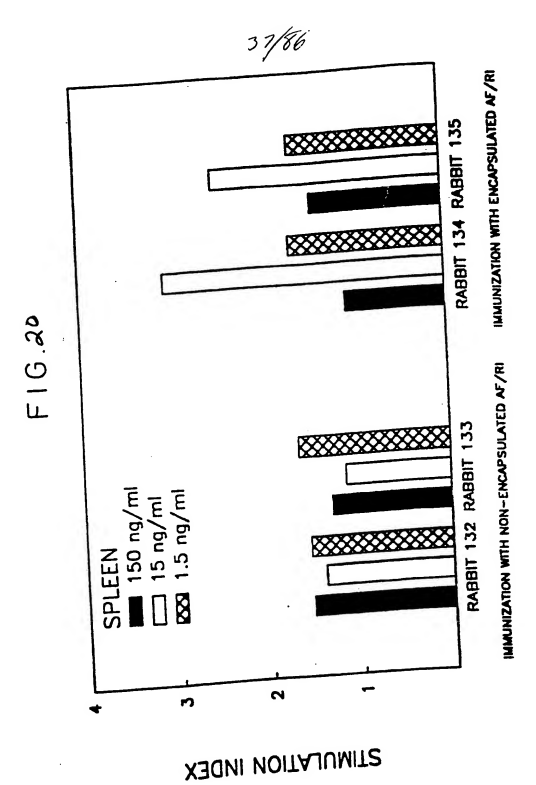


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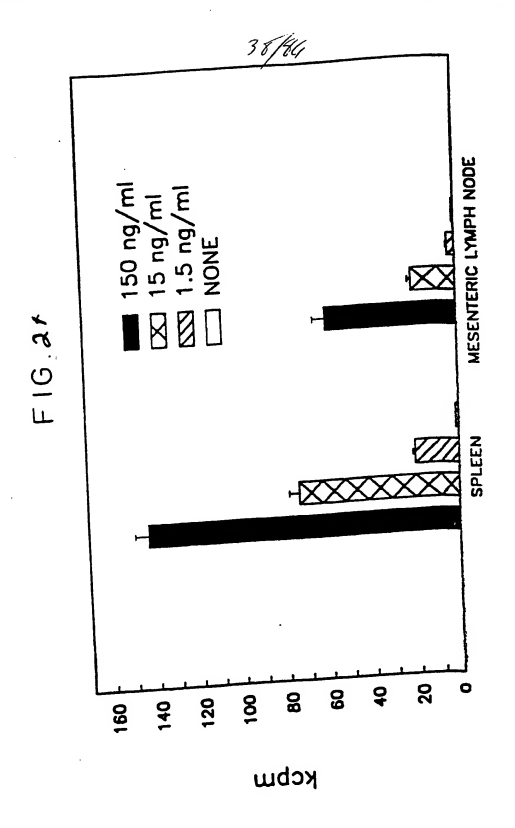


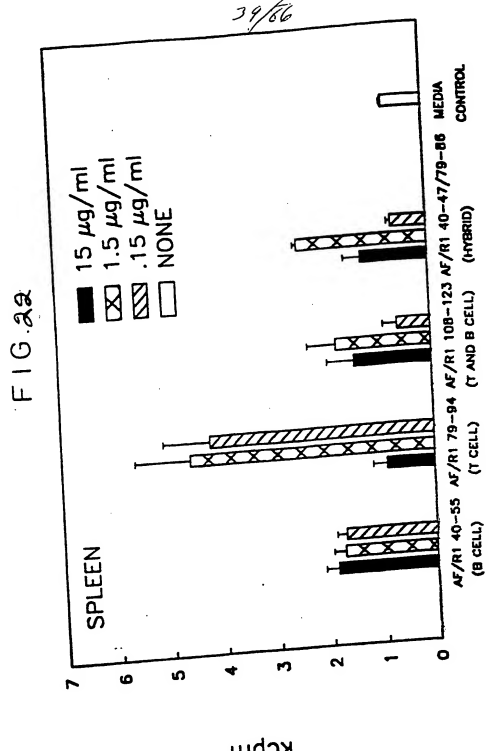


STIMULATION INDEX

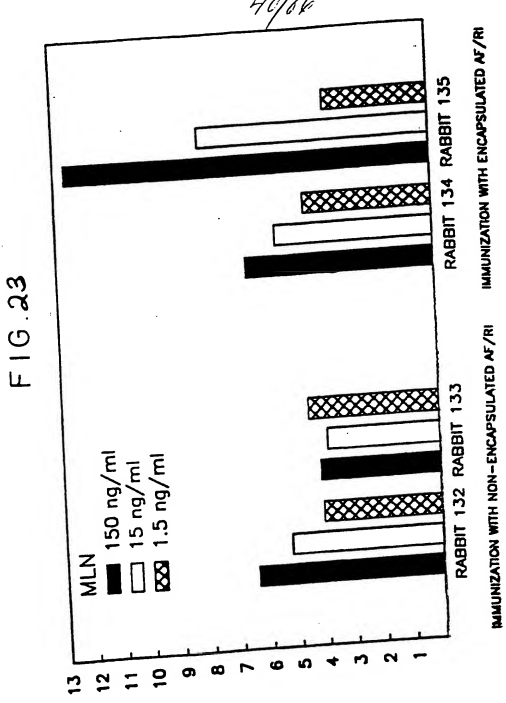


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4//8/ FIG 24 a 1 2 3

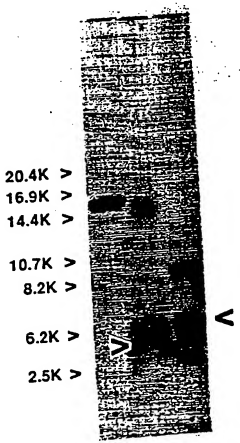
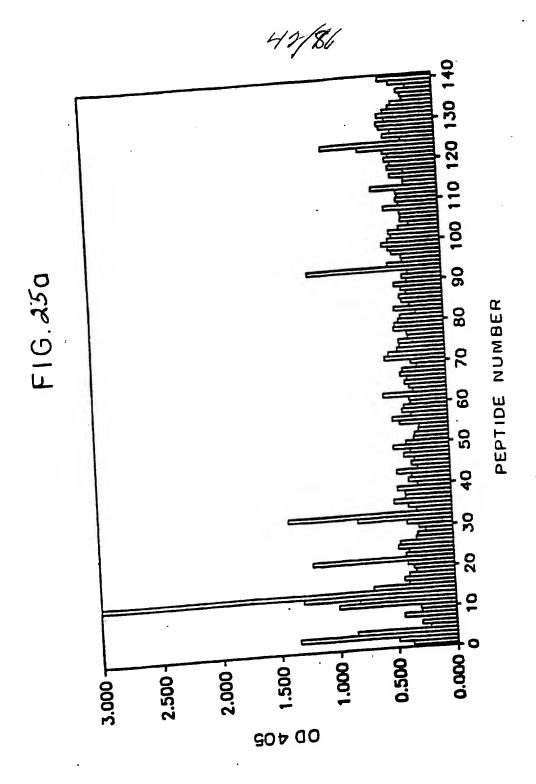
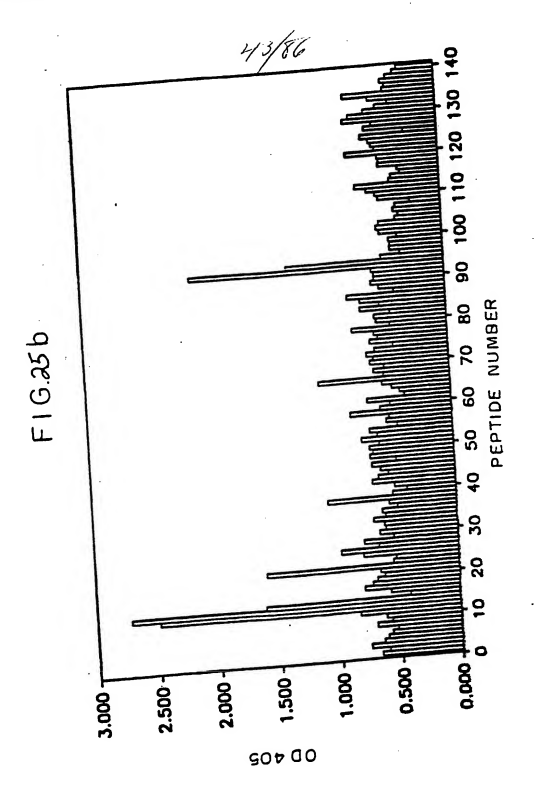
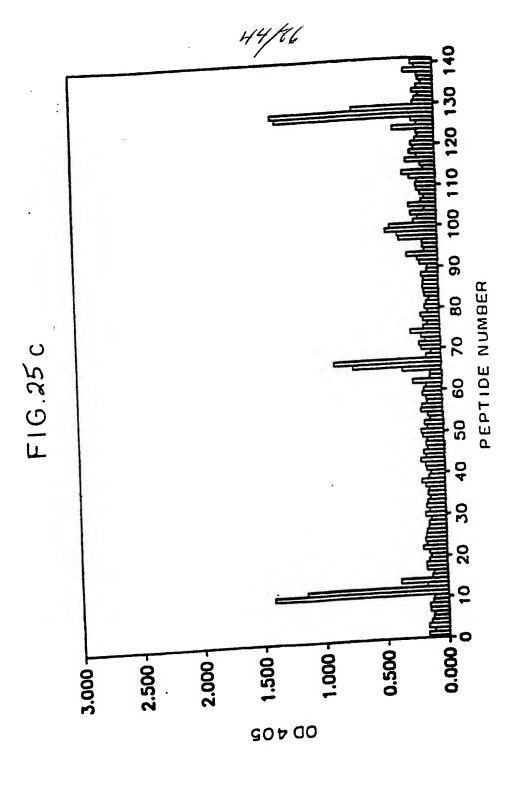


FIG.24b

Lane 2 LADTPQLTDVLNSTVQMP (62-79)
Lane 3 SYRVMTQVHTNDATKKVIV (42-60)

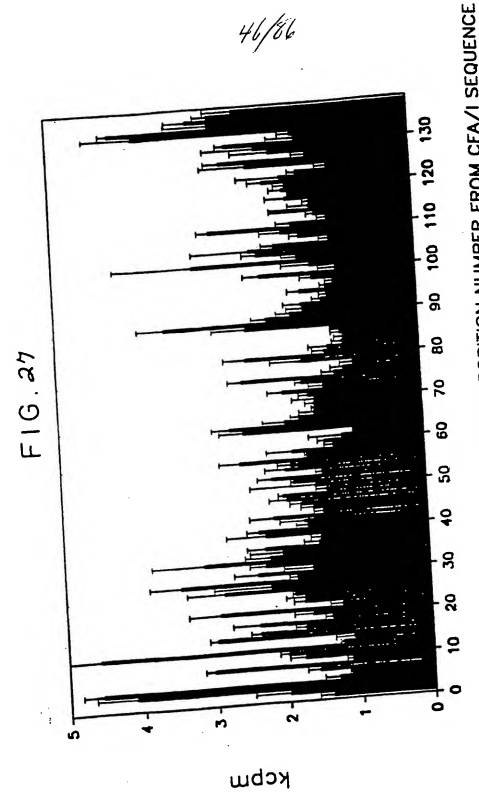






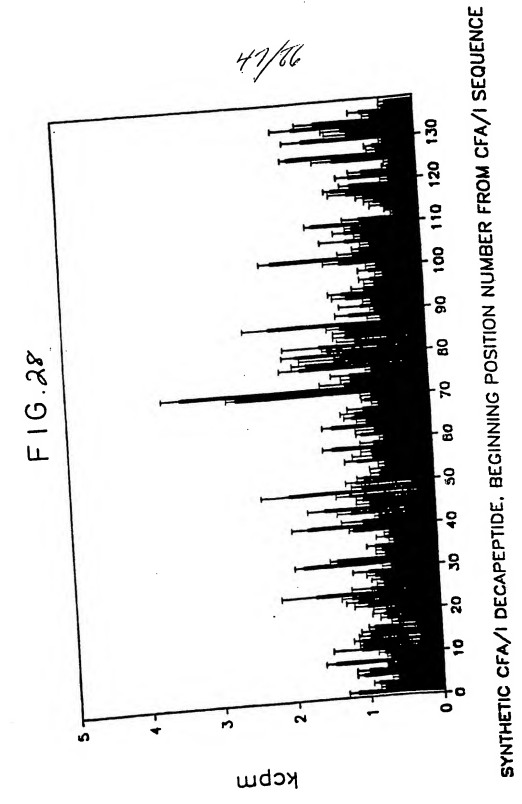
F16.26

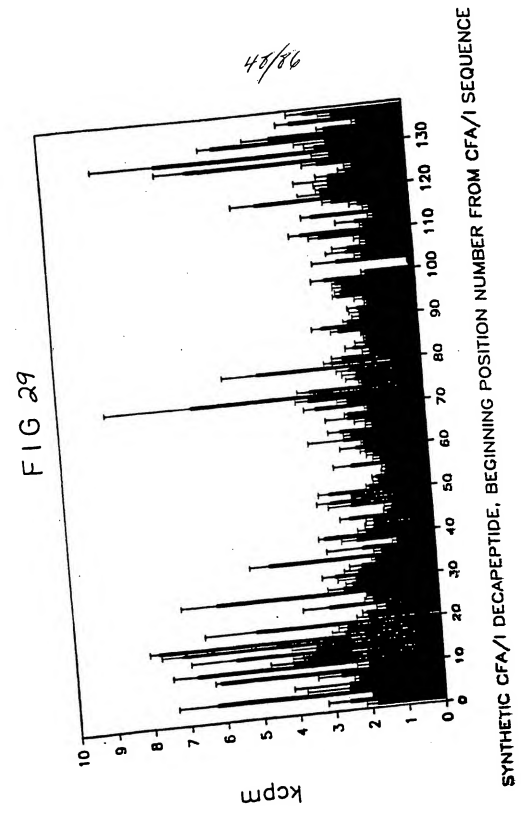
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TOLLQADGNALPSAVKLAYSPASKTFESYRVHTOVH IDLLQADGNALPSAVKLAYSPASKTFESYRVHTOVH IIDLLQADGNALPSAVKLAYSPASKTFESYRVHTOVH	TPOLTDVLNSTVQMPISVSMGGQVLSTTAKEFEAAA TPOLTDVLNSTVQMPISVSMGGQVLSTTAKEFEAAA TPOLTDVLNSTVQMPISVSMGGQVLSTTAKEFEAAA TPOLTDVLNSTVQMPISVSMGGQVLSTTAKEFEAAA	OTAGNYSGVVSLVMTLGS PTAGNYSGVVSLVMTLGS PTAGNYSGVVSLVMTLGS
		VSSSQELVISAAPKTAGTAPTAGNYSGVVSLVMT VSSSQELVISAAPKTAGTAPTAGNYSGVVSLVMT VSSSQELVISAAPKTAGTAFTAGNYSGVVSLVMT
VEKNITVTASUDPV VEKNITVTASUDPV	60. THDATKKVIVK THDATKKVIVK	TO Y S A S G V M G TO Y S A S A S G V M G TO Y S A S G V M G TO Y S A S G V M G TO Y M
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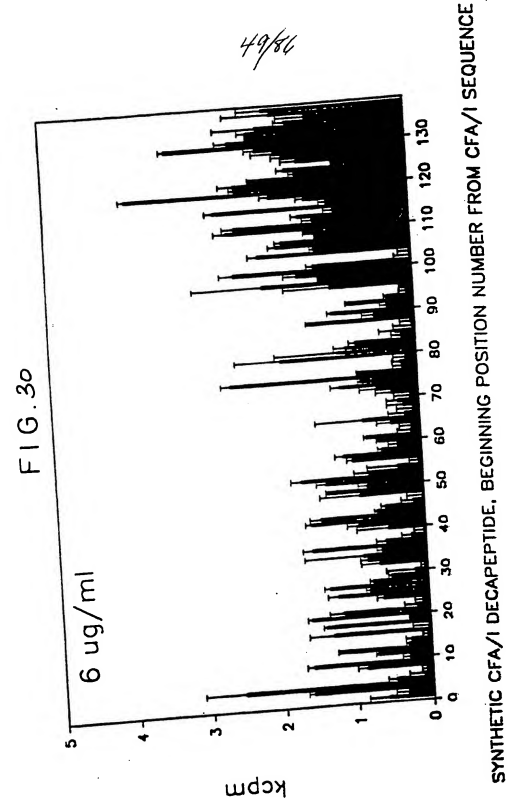
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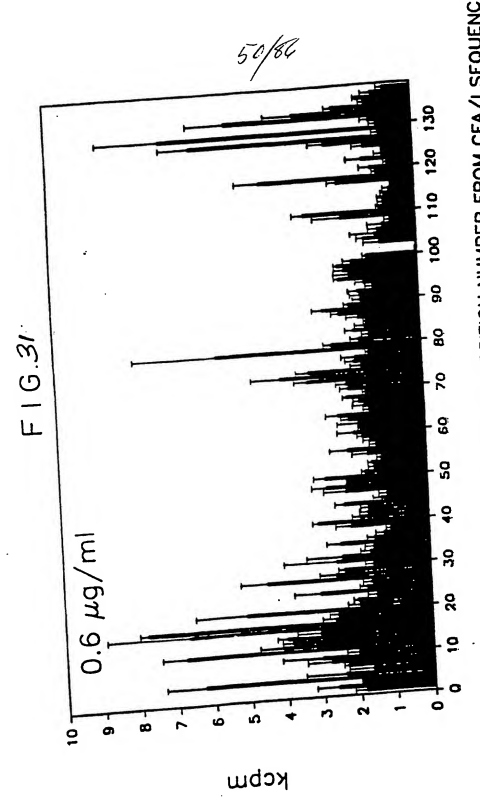
PCT/US98/01556 WO 98/32427





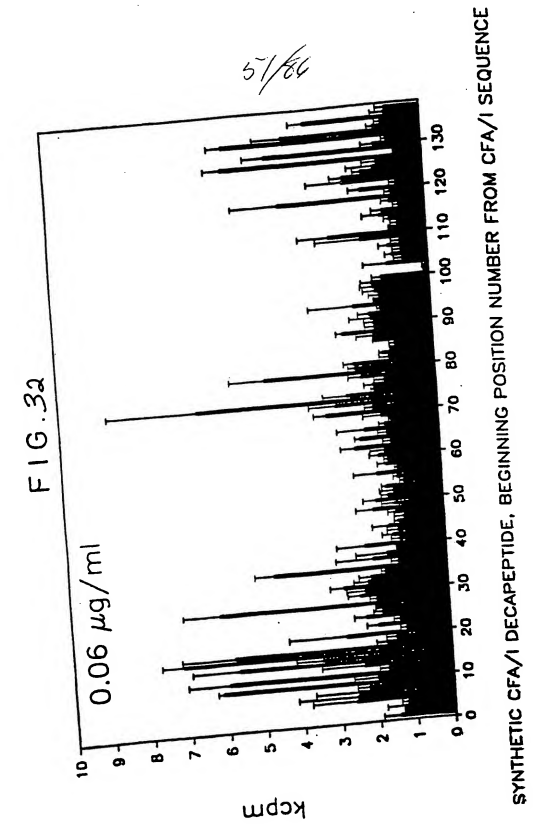
PCT/US98/01556 WO 98/32427

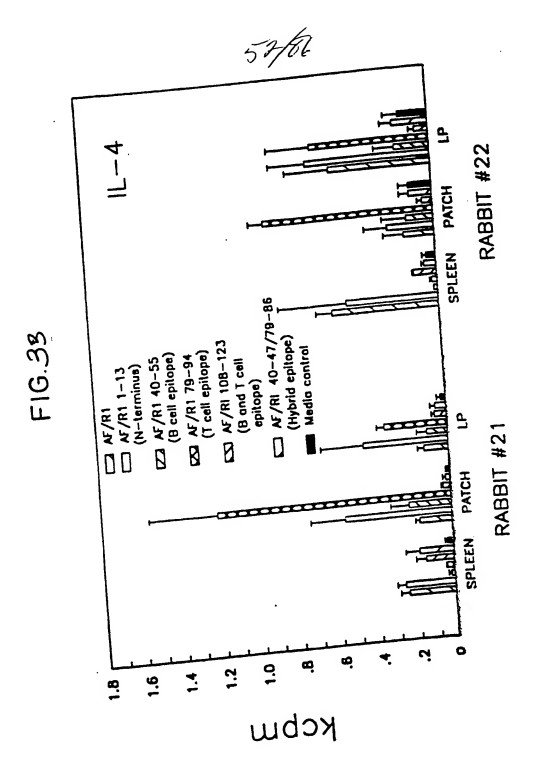


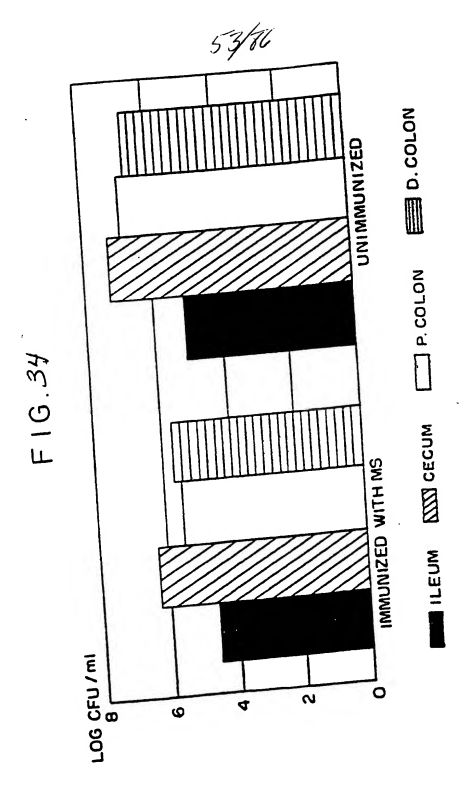


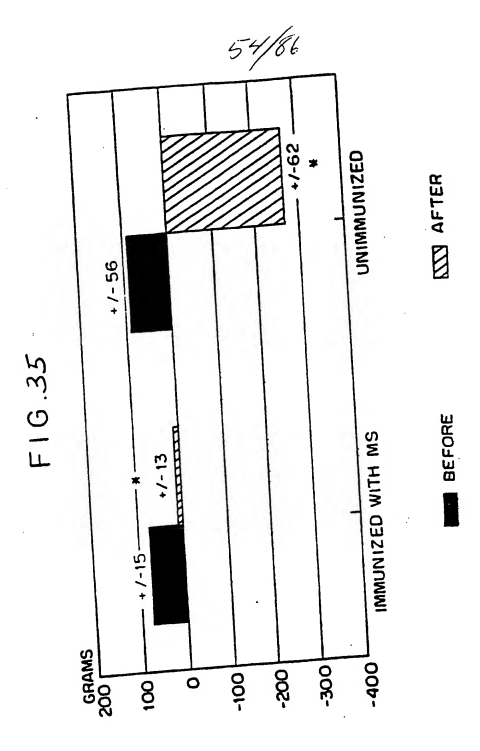
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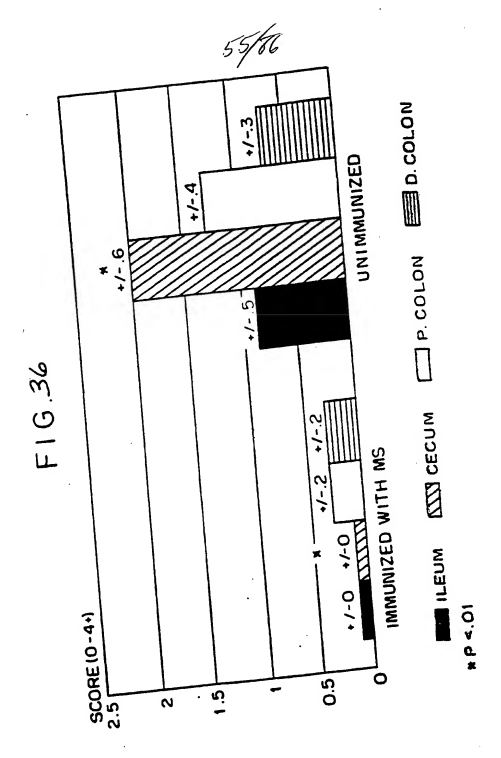
PCT/US98/01556 WO 98/32427

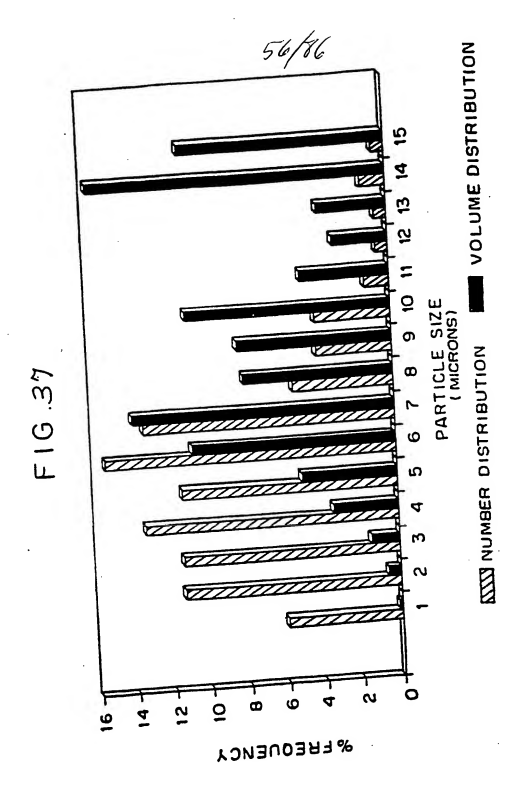






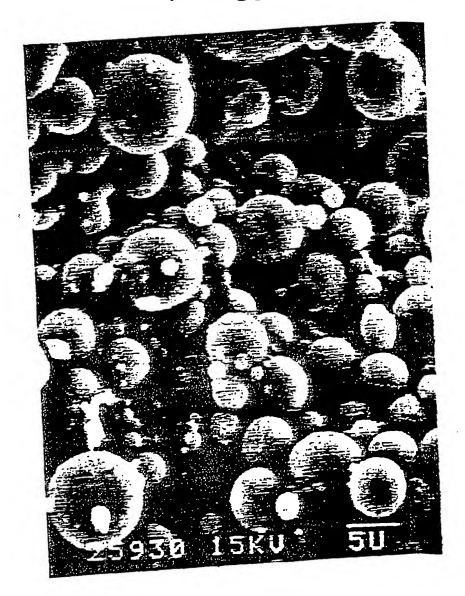


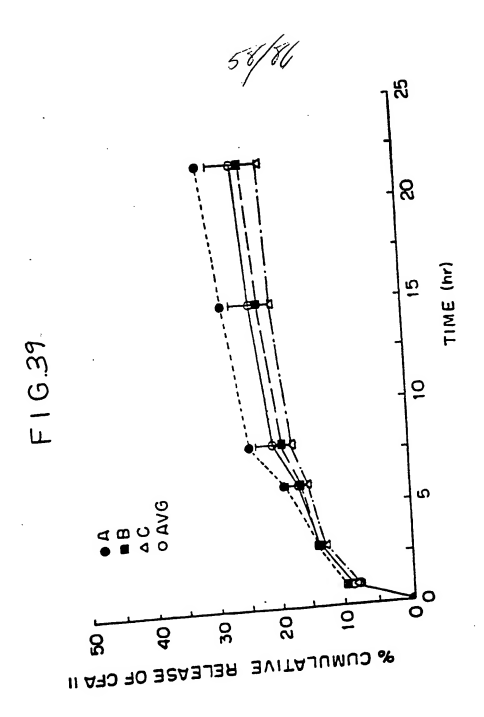


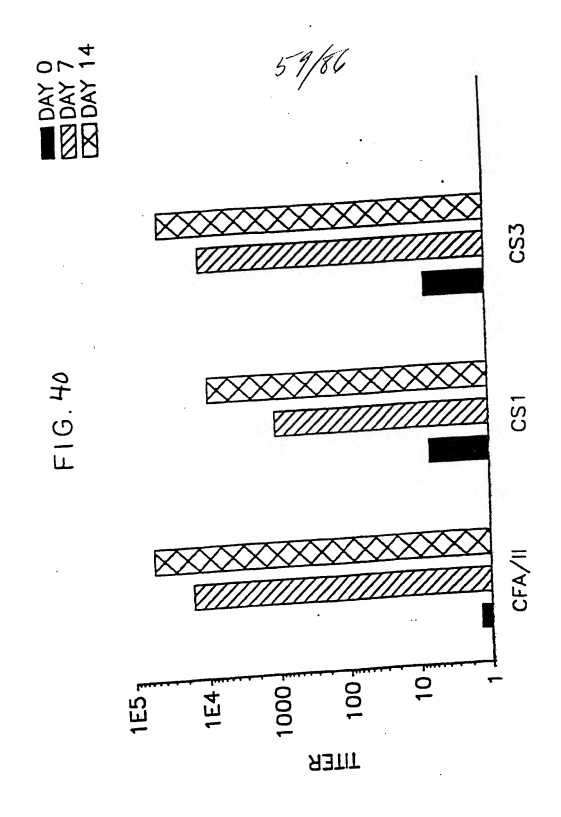


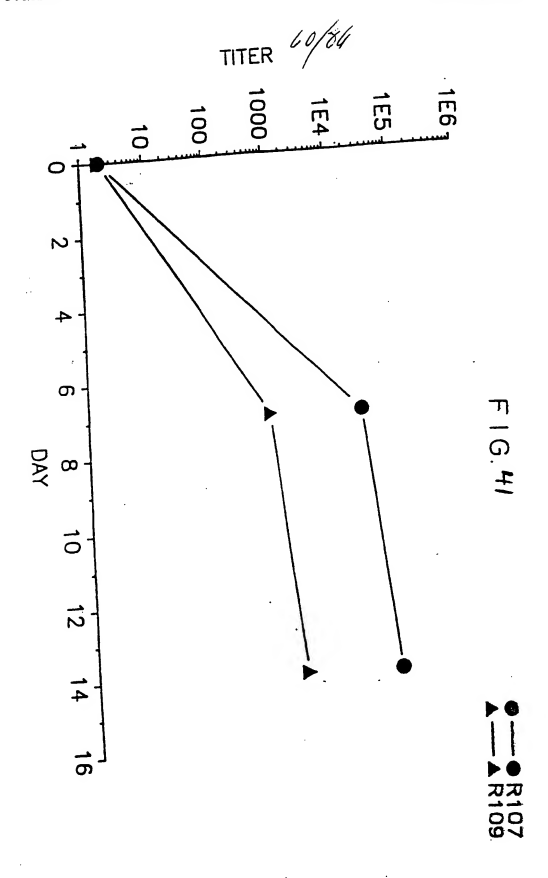
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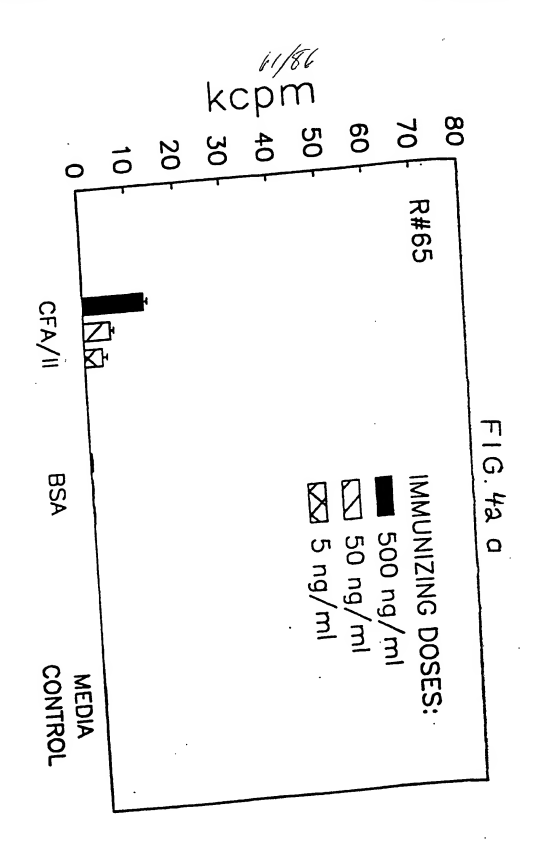
57/86 FIG 38

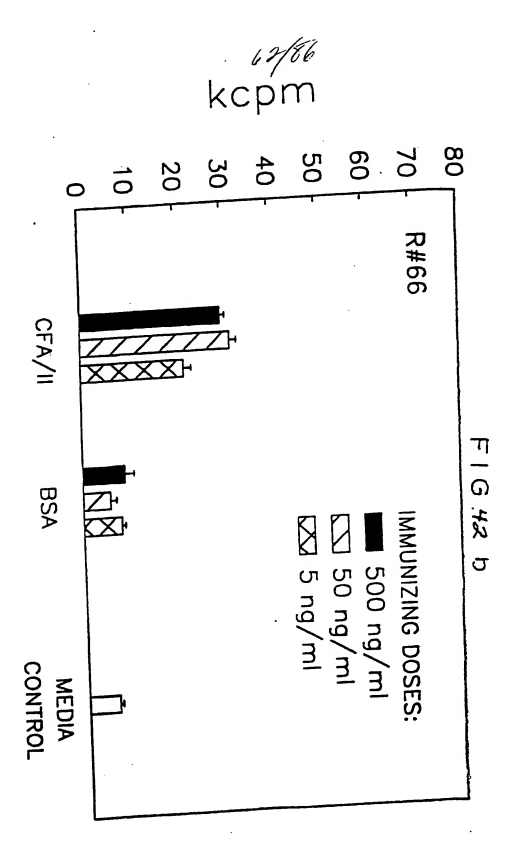


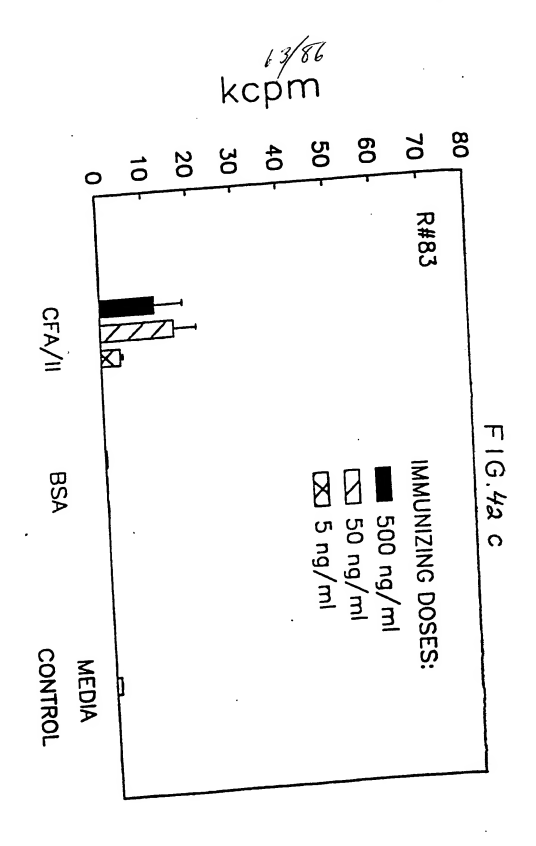


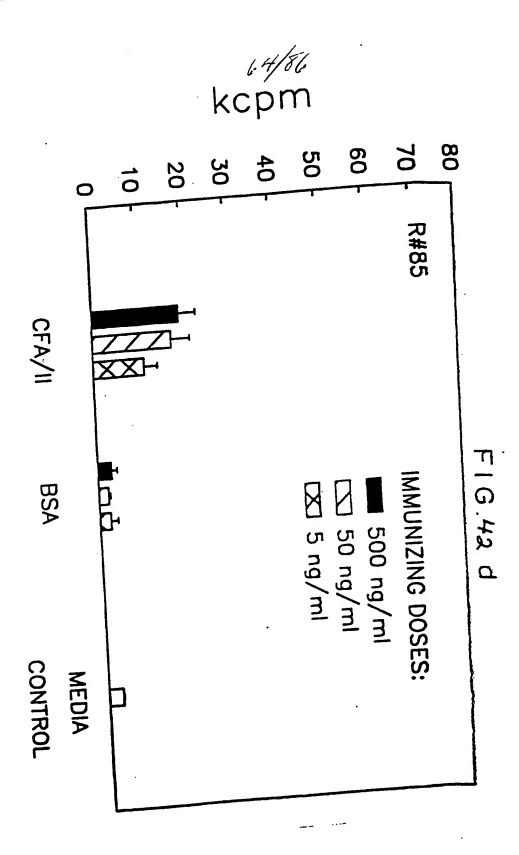


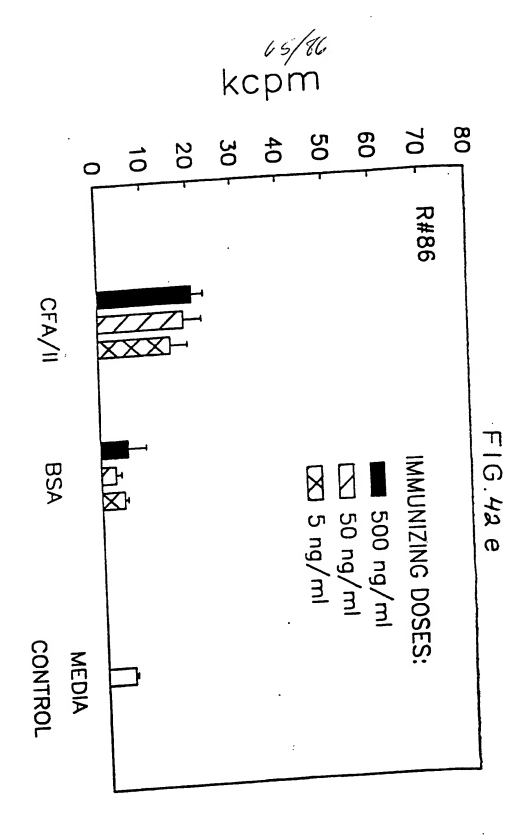


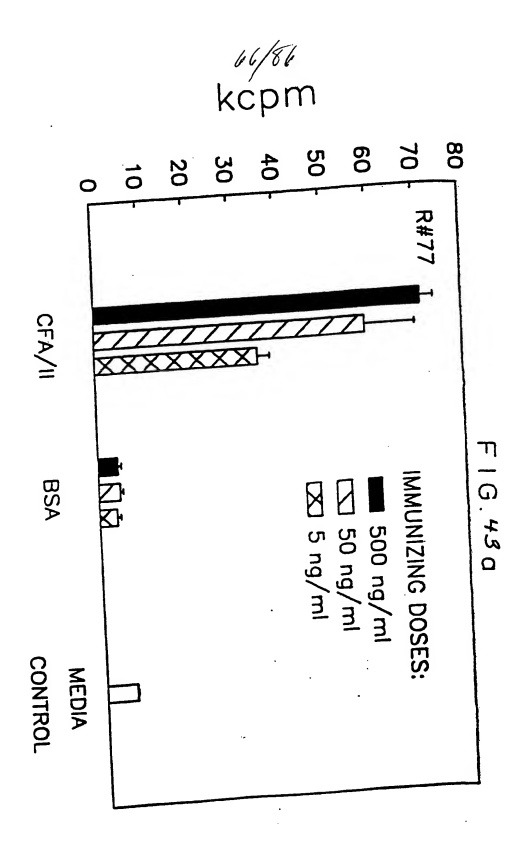


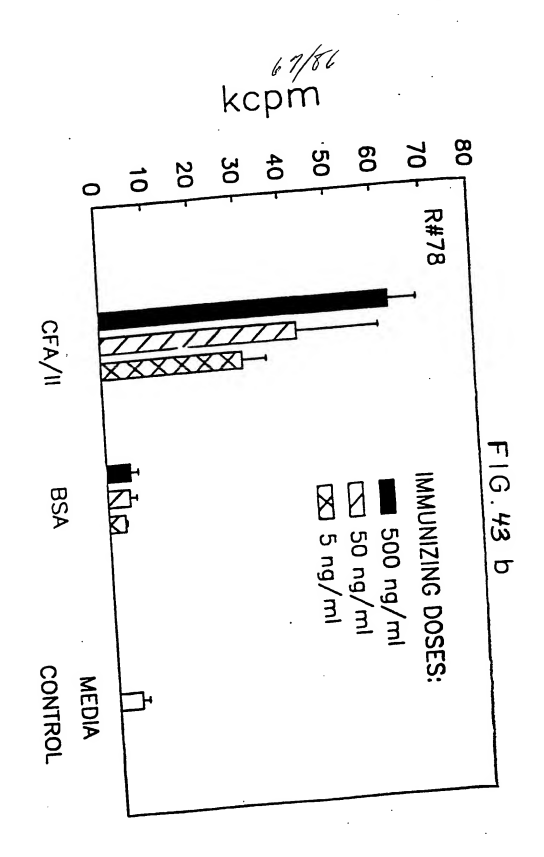


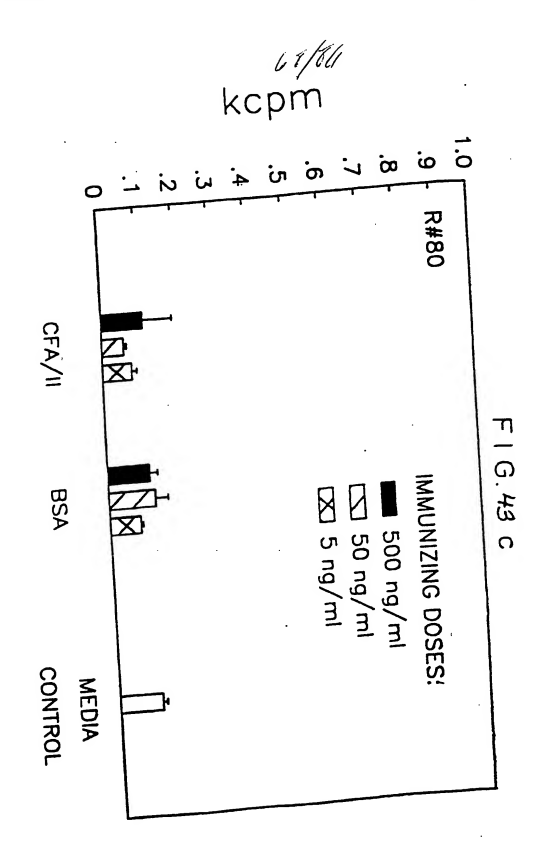


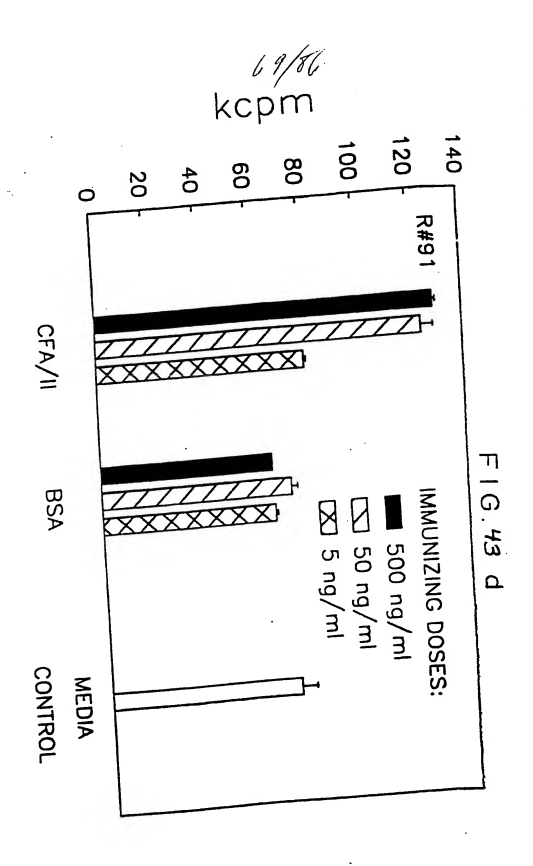


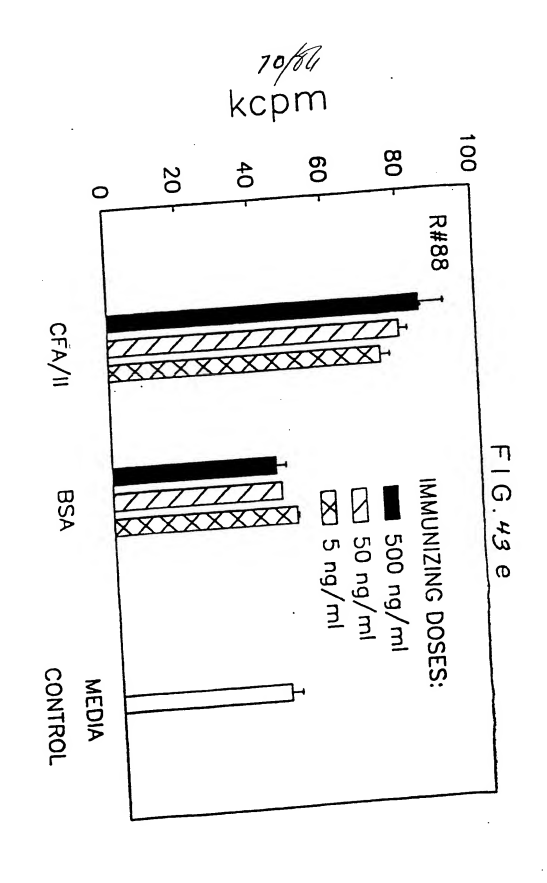


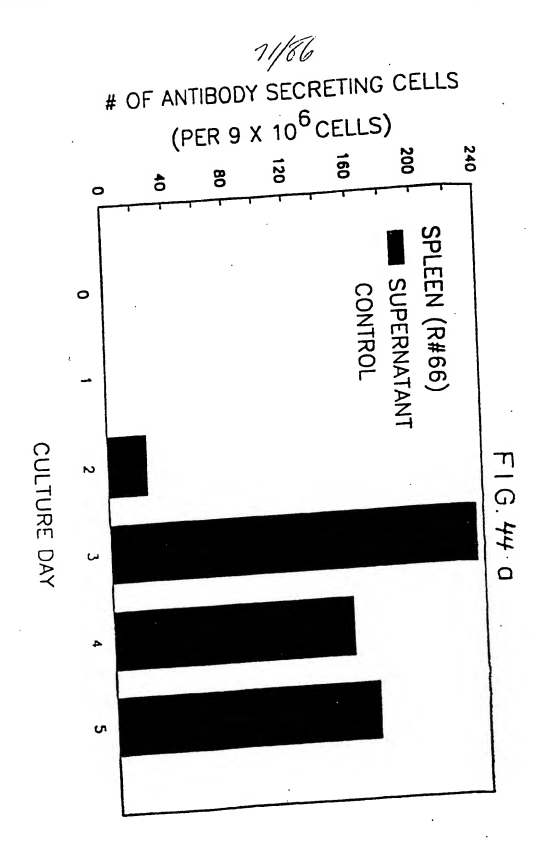


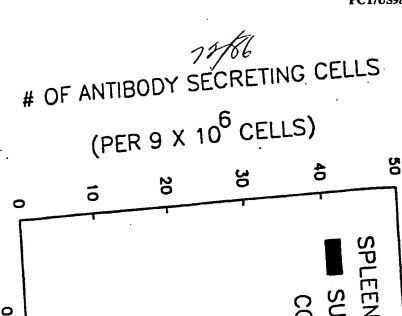


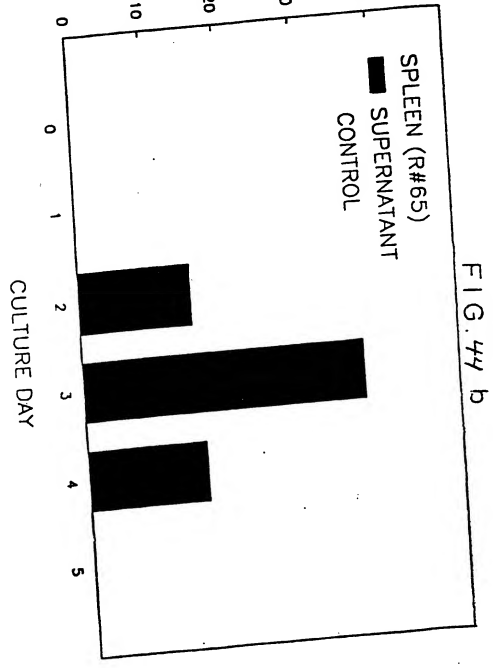


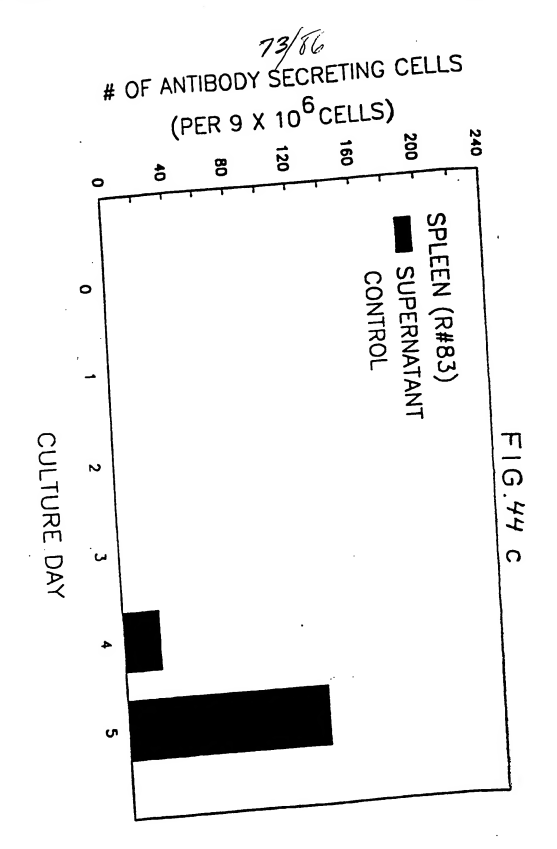


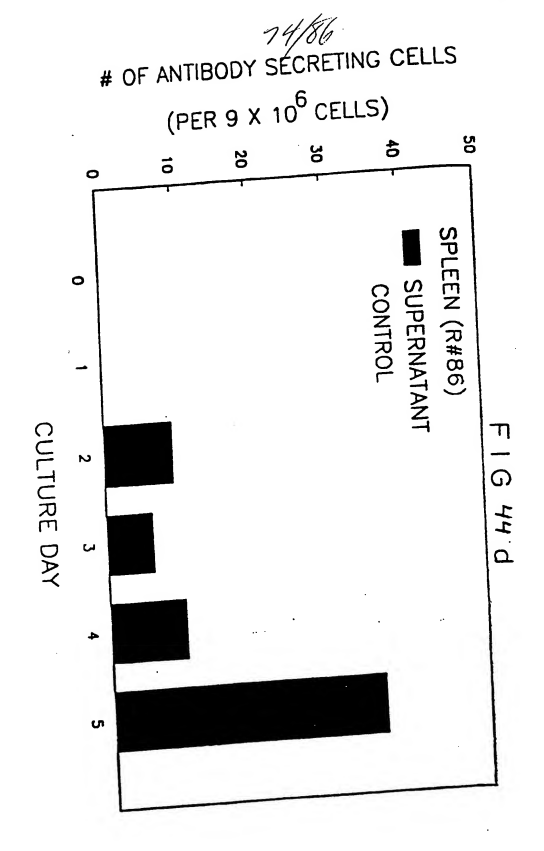


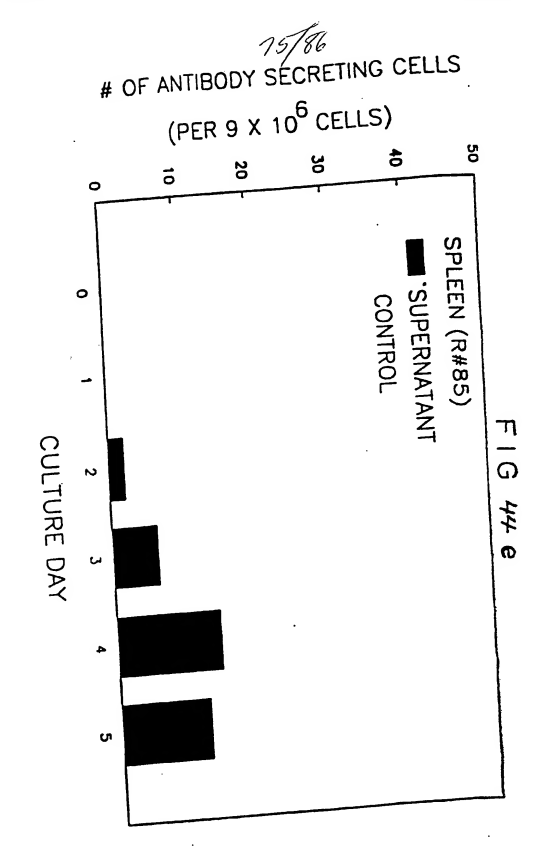


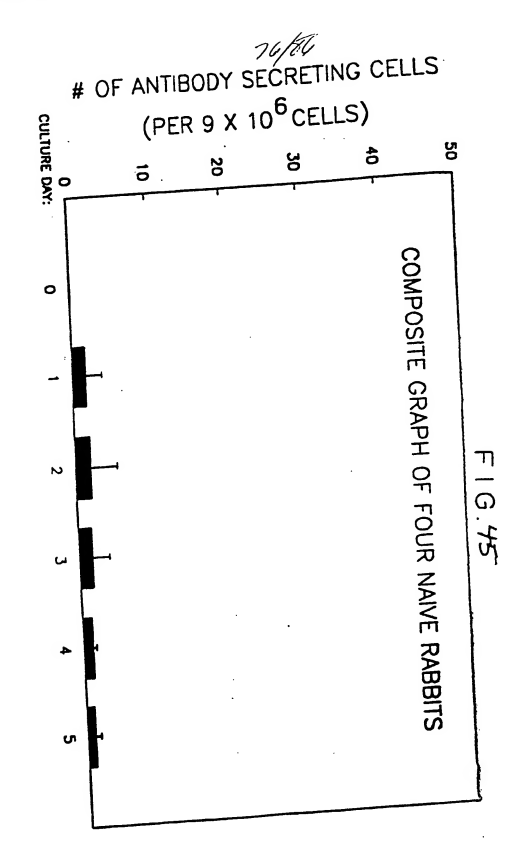




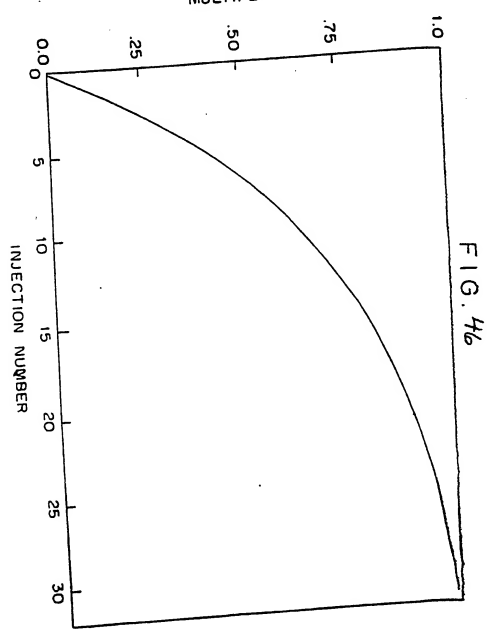


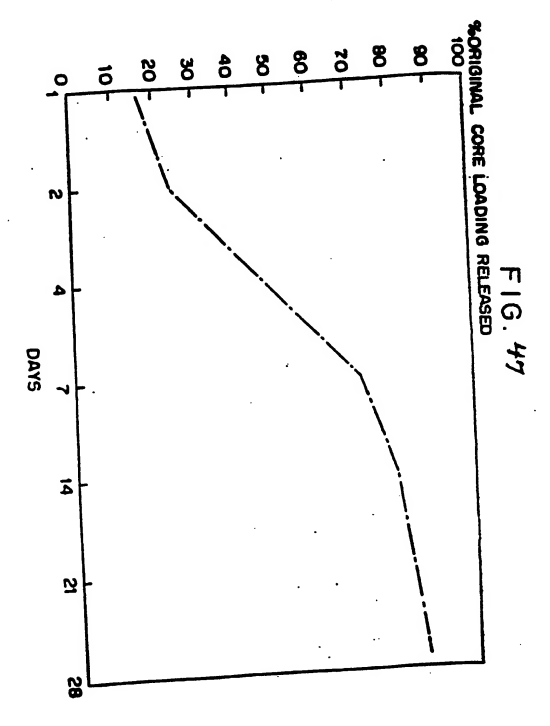






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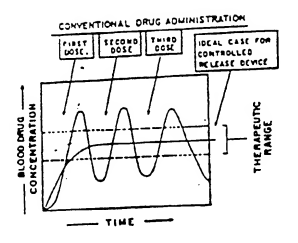


Figure 48

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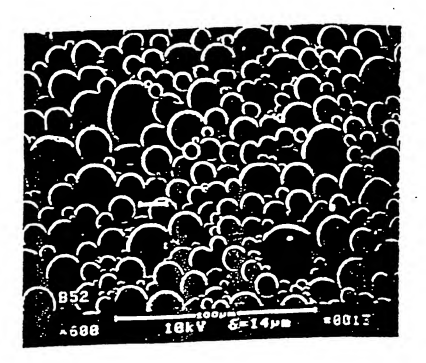


Figure 49

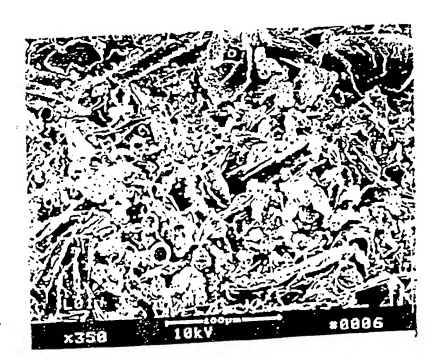
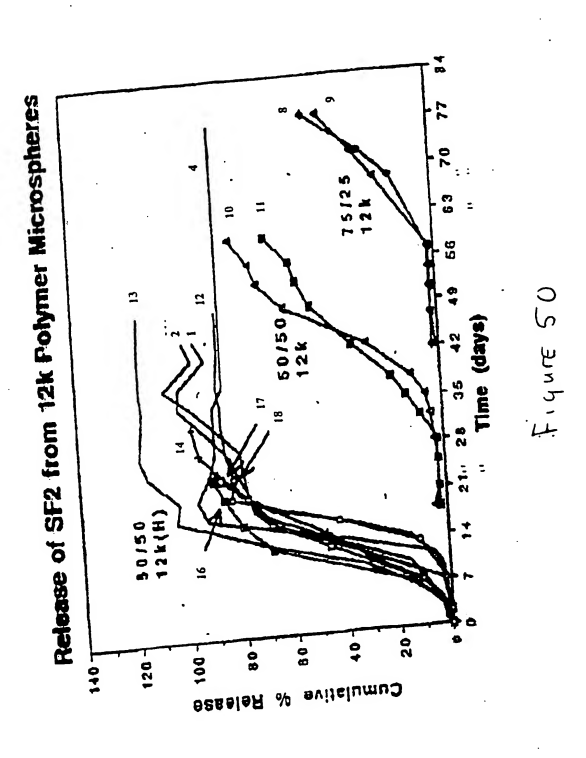


Figure 49a



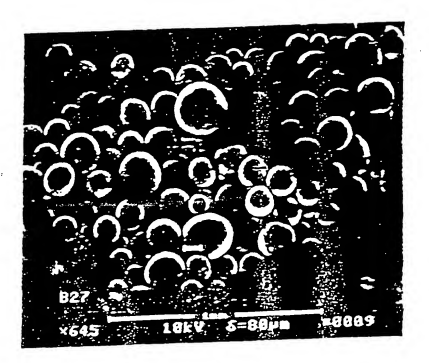
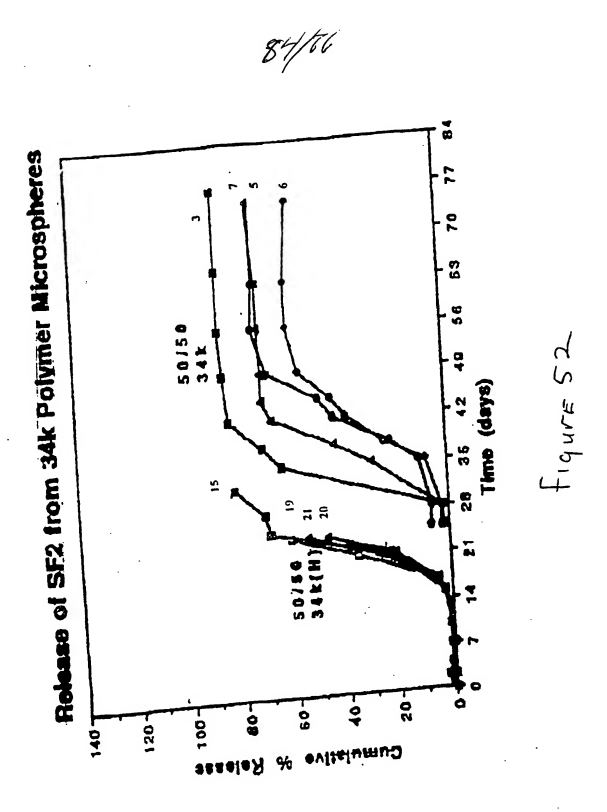


Figure 51



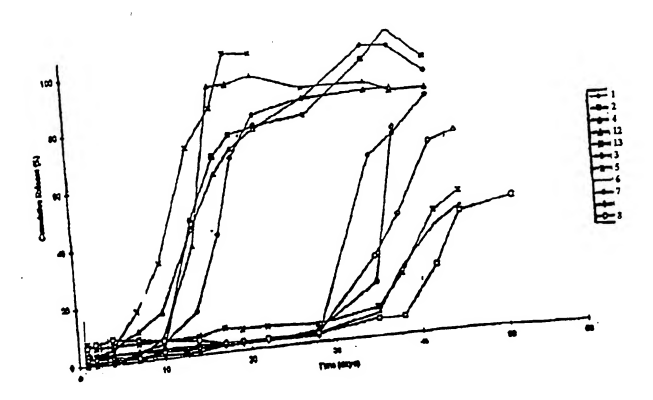
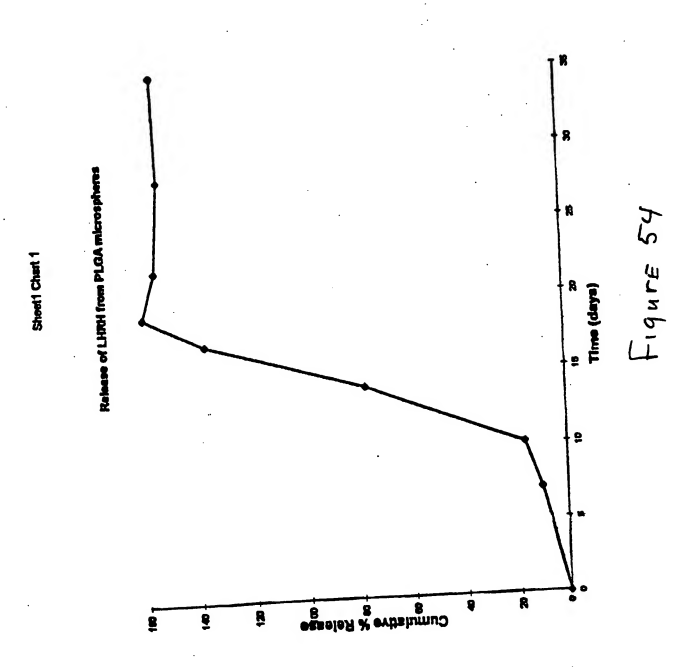


Figure 53





International application No. PCT/US98/01556

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :A61K 9/52; A61K 47/30							
US CL :424/501	US CL :424/501 According to International Patent Classification (IPC) or to both national classification and IPC						
B. FIELDS SEARCHED							
	ation searched (classification system followed	by classification symbols)					
U.S. : 424/426, 484, 501; 525/411, 413, 415; 528/354							
Documentation search	hed other than minimum documentation to the	extent that such documents are included	in the fields searched				
Electronic data base	consulted during the international search (na	me of data base and, where practicable,	search terms used)				
CAS ONLINE							
C. DOCUMENTS CONSIDERED TO BE RELEVANT							
Category* Cita	ation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.				
Micro	JEYANTHI et al. Novel, Burst-Free, Programmable Biodegradable Microspheres for Controlled Release of Polypeptides. In: Proceedings International Symposium on Controlled Release of Bioactive Materials 1996. Pages 351-352.		1-12, 21, 37, 42- 48, 155				
			13-18, 20, 32-36				
Conti	NG et al. Influence of Formulation rolled Release of Protein from Polonolled Release. September 1991,	42-48					
	0052510 B2 (SYNTEX (U.S.A.)I e document.	1-18, 20, 21, 32- 37, 42-48, 155					
N Susther dogue	ments are listed in the continuation of Box C	. See patent family annex.					
Special categories of cited documents: "T" later document published after the international filing date or priority data and not in conflict with the application but cited to understand the principle or theory underlying the invention							
L document wh	*B* earlier document published on or after the international filing date "X° document of particular relevance; the claimed invention cannot considered novel or cannot be considered to involve an inventive						
cited to establish the publication date of another citation or other special reason (as specified) *O* document referring to an oral disclosure, use, exhibition or other means **C* document referring to an oral disclosure, use, exhibition or other being obvious to a person skilled in		step when the document is h documents, such combination					
P document published prior to the international filing data but later than *&* document member of the same pater the priority data claimed			t family				
Date of the actual completion of the international search Date of mailing of the international search			arch report				
22 MAY 1998		18 JUN 1998					
Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231		ROBERT H. HARRISON (783) 308-2422	Jelingo				
Lacamina 140. (V	00) 000-0400	1010huoho 170(1(1. 000 2122	(/				

International application No.
PCT/US98/01556

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C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant	Relevant to claim No	
Y	YAN et al. Characterization and Morphological Analysis of Protein-loaded Poly(Lactide-co-Glycolide) Microparticles Prepared by Water-in-Oil-in-Water Emulsion Technique. J. of Controlled Release. 1994, Vol 32., No. 3., pages 231-241.		42-48
Y	YEH et al. A Novel Emulsification-Solvent Extraction Technique for Production of Protein Loaded Biodegradable Microparticles for Vaccine and Drug Delivery. 1995, Vol. 33, No. 3, pages 437-445.		42-48
Y	US 5,486,503 A (OPPENHEIM et al.) 23 January 1996 document.	5, see entire	1-18, 20, 21, 32- 37, 42-48, 155
	·		

International application No. PCT/US98/01556

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)				
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:				
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:				
Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:				
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).				
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)				
This International Searching Authority found multiple inventions in this international application, as follows:				
Pleaso Seo Extra Sheet.				
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.				
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.				
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:				
4. X No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-18, 20-22, 32-37, 42-48 and 155				
Remark on Protest The additional search fees were accompanied by the applicant's protest.				
No protest accompanied the payment of additional search fees.				

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BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In addition, this application contains claims directed to more than one species of the generic invention. These species are deemed to lack Unity of Invention because they are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for more than one species to be searched, the appropriate additional search fees must be paid.

<u>Oroup I(a)-I(d)</u>drawn to a composition for the burst-free, sustained, programmable release of active material(s) over a period from 1-100 days, process of using composition for human administration via parenteral route, process for the manufacture of the composition.

Groups I(a)-I(d) represent species of the genus "active material" which lack common structures and modes of action. Moreover, the recited Markush group of claim 8 contains redundant members; for example, peptides encompasses hormonal peptides as well as antibacterial peptides. Such extensive overlap prohibits a meaningful separation of the myriad inventions present with the exception of specifically recited species.

Group I(a) claims 1-18, 20-22, 32-35, 36, (37, 155) and 42-48 drawn to histatin containing compositions.

Group I(b) claims 1-10, 19-22, 31, 36, (37, 155) and 42-48 drawn to LHRH peptide containing compositions.

Group I(c) claims 1-12, 21-26, 36, (37,129, 155) 42-48 and 114 drawn to compositions comprising enterotoxigenic polypeptides.

Group I(d) claims 1-11, 36, 37, 42-48 (37, 129, 155) and 114 drawn to compositions containing Hepatitis B surface antigen.

Claim 155 is included with claim 37 to the extent that the recited methods overlap. It is noted that claim 155 fails to recite a positive method step.

Group II, claims 1 and 28-30 drawn to compositions for the burst-free, sustained, programmable release of active material(s)containing an additional oil phase.

Group III., claim 38, drawn to a process of using the compositions of Group I via topical route.

Group IV., claim 39, drawn to a process of using the compositions of Group I via oral route.

Group V., claim 40, drawn to a process of using the compositions of Group I via nasal, transdermal, rectal, and vaginal routes.

Group VI., claim 41, drawn to a process of using the compositions of Group I via oral or nasal inhalation.

Group VII., claims 49-58, 100 and 101, drawn to a process for the protection against infection of a mammal wherein a peptide is employed as the "active material".

Group VIII(subgroups (1)-(29), claims (59-66, 102-111, 112-117, 118-120, 121-123 and 156) drawn to a vaccine compositions containing the polymers of claim 1 and one of the specifically recited species present in claims 70-92, (subgroups (1)-(23)) and 94-99 (subgroups(24)-(29)). Claims 67-69 and 93 will be searched with the appropriate specific peptide.

Group IX., claim 124, drawn to an assay employing composition for the burst-free, sustained, programmable release of active material(s).

Group X, claims 125-132, drawn to methods of preparing antibodies employing the compositions as described in claim 52.

Group XI., claims 134-154, drawn to a process for the protection against or therapeutic treatment of bacterial infection in the soft tissue or bone of a mammal wherein an antibiotic is employed.

The inventions listed as Groups I-XI do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: The generic



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composition set forth in claim 1 is taught by Jeyanthi et al.(1996) and therefore cannot constitute a special technical feature. In setting forth Groups I(a)-I(d) the unifying technical feature is viewed as the specifically recited molecule in combination with the polymeric composition of claim 1(b). Each of these specifically recited molecules differs from the others both in structure and mode of action and therefore represent distinct technical features. The additional uses recited in groups II-XI and subgroups therein are distinct and different in the overall manner in which each is carried out.						
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Form PCT/ISA/210 (extra sheet)(July 1992)*

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